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The Practices of Childhood
Coproducing Child Health in Aotearoa New Zealand

Julie Sera Spray

This thesis addresses the question: How do children participate in the coproduction of their health? Aotearoa New Zealand childhoods are characterised by persistent inequities which concentrate in the intersection of class and ethnicity. One manifestation of these inequities is the disproportionate burden of acute and chronic illness for Māori and Pasifika children, particularly those living in low socio-economic areas. A large, interdisciplinary body of research has considered questions of child health inequities from broader epidemiological and social perspectives, identifying the role of structural determinants of health: for example, links between housing and distribution of morbidity. Yet, as a robust anthropological literature on children’s agency demonstrates, children are not passive recipients of health care, but actively negotiate social relations and collectively produce cultural meanings and practices. What, then, are the impacts of such activities on child health; what are the processes through which children help to shape their own wellbeing?

I approach these questions through ethnographic fieldwork in a school with children aged between 8-12, located in an area of South Auckland characterised by social marginalisation and material deprivation. Drawing on the work of Corsaro, Bourdieu, Shilling, and Prout on relationships between individuals—or individual bodies—and society, I propose a coproduction framework which places children as participants in their health by positioning their activities in dialectical relationship with adults, the body, and social structures including the state, the institution, cultural ideologies, and the economy. Using this framework, I document how Tūrama School children’s practices are produced from their experiences of their bodies and these wider structures, and how these practices, such as monitoring the body, eating, using pharmaceuticals, negotiating health care, risk management, and constructing relationships can, in turn, help to pattern children’s bodies and health in significant ways. Findings from this thesis suggest ways to situate children’s embodied practices in relation to political-economic and social forces, elevating their perspectives and activities without conferring upon them sole responsibility for their own health status.
For Janice,
who loved these children
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Glossary

Note on conventions

Although common convention is to italicise non-English words, I have elected to forgo this convention for Māori words in this thesis (words in other non-English languages are still italicised). My reasons for this are twofold: firstly, Te Reo Māori is an official language of New Zealand, but one that is typically marginalised in favour of the dominant English. I therefore wish to avoid further ‘othering’ of Te Reo. Secondly, this format better reflects the way that Māori words are organically woven into every day written and spoken New Zealand English, particularly in my study community.

Some Māori words are not easily translatable into English. So as not to disrupt sentence clarity, some words requiring more detailed explanations are footnoted. Māori translations are referenced from the Te Aka Māori-English Dictionary (Moorfield 2011).

Where children have anglicised Māori words, for example by pluralising words by adding ’s’, I have represented the hybridised word as they have used it in context.

All words are Māori unless specified otherwise.

_āiga (Samoan):_ Wider family
_Aotearoa:_ New Zealand
_hapū:_ sub-tribe/clan
_hui:_ gathering, meeting
_iwi:_ tribe
_kai:_ food, eating
_kainga (Tongan):_ Extended family
_kaitiaki:_ guardian
_kapa haka:_ Māori cultural performance
_kaumātua:_ elder
_kōrero:_ speech, address
_marae:_ A central gathering place for Māori, technically meaning the courtyard in from of the whare nui (main meeting house) but often referring the key buildings of the complex.
_Pākehā:_ New Zealander of European ancestry (See chapter two for further discussion)
_pōhara:_ poor, impoverished
_poi:_ light balls on the end of strings that are rhythmically swung to accompany waiata
_tangihanga, tangi:_ Māori funeral, rites for the dead, weeping
_tonga:_ treasure
_Te Ao Māori:_ The Māori world
_te Ao Wairua:_ The spiritual world
_Te Reo:_ The (Māori) language
_tikanga:_ correct procedure, custom
_tūrama:_ light, to give light to, illuminate
_waiata:_ song, to sing
_wairua:_ spirit, soul
_whaea:_ Mother or aunty
whānau: extended family, family group (see footnote 3 of Preface for further detail)
whāngai: adopted (usually informally) child
whare nui: meeting house, main building of a marae, where guests are accommodated and hui (meetings) held
List of abbreviations

BPS: Better Public Services [targets]
CRHD: Chronic Rheumatic Heart Disease
CYF: Child, Youth and Family
DHB: District Health Board
DSM-V: Diagnostic and Statistical Manual of Mental Disorders, 5th edition
GAS: Group A Streptococcus [bacteria]
IQ: Intelligence Quota
MP: Member of Parliament
NGO: Non-Government Organisation
NSSC: New Social Studies of Childhood
NSSI/NSSID: Non-Suicidal Self Injury/Disorder
PTSD: Post-Traumatic Stress Disorder
RFPP: Rheumatic Fever Prevention Programme
WHO: World Health Organisation
Chapter One

Introduction

Nine-year-old Victor ‘cannot tell a lie’, his mother Adrienne tells me. The youngest of four boys, he would go out with his brothers, who would warn him, ‘don’t tell Mum that we went for a kai [feed].’ Back home, Adrienne would ask if he enjoyed his time. ‘Yup,’ Victor would reply, ‘I didn’t go to McDonalds.’

‘Oh you went to McDonalds, aye?!’

‘No I didn’t!’ Victor would protest. ‘I didn’t have a cheeseburger!’

Victor’s honesty, which could embarrass his mother and frequently got him into trouble at school, was much more useful to me as a researcher. Making lists at his kitchen table about ‘what it means to be a child’ and ‘what it means to be an adult’, Victor’s mother puts her head in her hands as he explains to me that ‘adults don’t have enough money, so you got to drip dry your towels’, that adults had to put their cold food in the cupboard rather than a refrigerator, and that adults ‘need money to afford a lighter, or you can’t buy your packets of smokes’.¹

It seems that Victor’s experience of adults is seeing them struggle. His ideas about childhood and adulthood are shaped by the material deprivation of his family, in turn a product of national and global political, economic and historical forces. A history of colonialism and neoliberal economic policy in Aotearoa², or New Zealand has widened inequality and exacerbated a housing crisis within the Auckland region, both of which disproportionately affects Māori families like Victor’s, as well as many of his Pasifika classmates (Amore 2016). For those growing up in poverty, these conditions of inadequate housing, nutrition, and health care in turn create a particular experience of childhood. As this thesis will show, how children interpret the circumstances of their childhoods matters, not least because these understandings shape their practices: the way they negotiate their relationships, the way they

¹ Children’s speech from interviews, transcribed verbatim, has been lightly edited for clarity. I have not altered their grammar. Interactions outside of formal interviews were not recorded and so appear as reconstructions.
² Aotearoa, commonly translated as ‘land of the long white cloud’, is the Māori name for New Zealand and its use is widespread in New Zealand. The original derivation is unknown, but it appears to have been standardised in part through its inclusion in the 1878 Te Reo Māori translation of the New Zealand National Anthem, God Defend New Zealand, by Judge Thomas H. Smith of the Native Land Court.
approach risk, and the way they manage their bodies. While the political and economic forces of society shape their health from above, this is not the whole story of children’s health, rather, children’s interpretations of and responses to their experiences further mediate these effects. Child health is therefore a coproduction, and a full understanding of the dynamics of child health requires both a view of the broader workings of society, and a view from children’s eyes, as experienced from within a child’s body.

Victor’s experience of childhood, like many other children I worked with, is one of chronic ill health originating from the quicksand of poverty. When Victor was four, Adrienne had moved into a three-bedroom house owned by an acquaintance. The family soon discovered the house was in poor condition. ‘We were in a house that kind of lifted up a bit off the ground,’ Adrienne explains, ‘and underneath the house was full of water. It was full of water, and it wasn’t insulated-

‘-There was holes in the roof,’ Victor interjects.

Six months after they moved in, five-year-old Victor was hospitalised with heart valve damage following rheumatic fever.

From Adrienne’s perspective, the poor housing conditions were to blame for Victor’s illness. He was healthy before they moved. Adrienne tried asking the landlord for heating, and finally tried taking the case to court, without success. With a poor credit rating and supporting four children on a single income, Adrienne found herself trapped; while she had a roof over her head, Housing New Zealand would not assess her for State housing, but her landlord would not make the improvements she needed. The family remained in the house for another four and a half years, while Adrienne researched the causes of rheumatic fever and worked to combat the dampness that the landlord would not address. But the house was sitting atop about ten centimetres of water, while the surrounding grass would sink after rain to form a pool so deep ‘you could have like a swim there.’ Adrienne scrubbed the mildew-riddled rooms every week so that her children could breathe.

After five years, the rent was raised so high that Adrienne could no longer afford to pay. The family moved in with Adrienne’s sister—where I first visited them—and applied to Housing New Zealand [the State housing provider] for assistance. Due to Victor’s heart condition, the family were prioritised and when I visited them again three months later, they had relocated

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3 Government-owned and managed low-cost rental housing
to a State house in the same area. Adrienne was delighted with her new home: insulated, clean and dry.

Yet Victor’s body will always bear the marks of the conditions of his early life, and this is the body through which he experiences his childhood. At nine years old, he understands his heart has been damaged and he cannot play rugby league like his brothers and classmates, or go on the FearFall ride at the local theme park. Between his heart condition and his asthma he struggles to move physically, and, along with poor nutrition, as a consequence is obese. His large size has likely contributed to his sleep apnea, which keeps him awake at night, as does his eczema, but even more so his coughing from asthma. He sleepwalks, and his mother often finds him in front of the television in the wee hours of the morning when he can’t sleep. His skin is riddled with rashes, pock-marks and small sores, his nose runs from hayfever and he is prone to nosebleeds. Some of these conditions may improve, but much of this has set his life along a trajectory of poor health and social stigma.

How do children participate in the coproduction of their health?

Victor’s heart condition is currently managed under the District Health Board (DHB) with monthly prophylactic penicillin injections, administered at his school by a visiting public health nurse. Victor is not a passive recipient of care, however, but responds with his own set of practices, borne from his social environment, including at school. When he is called to the office to get his injection, he tells his class he is in trouble, reinforcing his identity as social deviant. He steals packets of biscuits from Adrienne’s cupboard to take to school for lunch, which, whether he intends or not, inevitably end up shared amongst the children in his class. On one day, unable to swallow the antibiotics administered by the school clinic for a strep throat, he covertly spat his pills into the bin, covered them with paper, thanked the nurse and returned to class. His health, therefore, is a product not only of what is done to him, but of how he makes sense of and negotiates a body lived within a social world he himself helps to create.

The way that children understand and respond to illness and its treatment therefore can have significant impact on how policies play out and what consequences they might have. The structural determinants of child health in Aotearoa have been extensively discussed in health

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*4 Governing body responsible for funding and providing health care to the local region.*
and social science literature, and are, to some extent, addressed in public health policy and services which, for example, provide subsidised insulation to improve housing, or targeted funding for community services (e.g. Telfar-Barnard et al. 2011). While the roles of the state, medical professionals, schools and parents are accounted for within child health policy, the roles children themselves play in their own health care tend to remain invisible. With a view of children as passive, lacking competency and vulnerable, the public policy approach to children since the landmark 1989 Children, Young Persons and Their Families Act has increasingly been oriented towards their protection via the regulation of their lives (Tap 2007) within a social investment paradigm concerned with maximising human capital and minimising future expense (Elizabeth and Larner 2009; Keddell 2018; O’Bien 2016). This assumption of adult control over children’s lives may underlie the lack of attention paid to children’s experience of health policy and of the body itself. But what happens to children’s experience of their bodies when they are asked every day to check on an aspect of their health? How do children come to define ‘lunch’ when governments say that good parents provide them with meals they do not have? What happens to children’s health care practices when illness is discursively linked to death?

These are not only questions about individual behaviour, but about how broader structural conditions (institutional, socio-economic, political, demographic) are experienced through bodies and translated into social meanings and practices within children’s peer cultures. I argue that the answers to these questions matter because the social meanings children produce from their structured, embodied experience, in turn reinforce, modify, or generate biosocial practices of the body.

I use the term ‘coproduction’ to characterise these processes, though what I mean by coproduction varies slightly from how other scholars have used the concept. From a Science, Technology and Society (STS) perspective, Sheila Jasanoff (2004) refers to the mutually constituting nature of science and society, whereby social, legal and political institutions both shape, and are shaped by scientific knowledge and associated technologies. Here, the coproduction framework powerfully reveals how science, while purporting objectivity, becomes imbued with the biases and political motives of the humans who create it, as well as how, in part because of this presumed objectivity, science and technology can modify or legitimise the institutions, power structures and values of society. Meanwhile, Margaret Lock

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5 In July 2017, as part of a suite of changes, the Act name was changed to the Oranga Tamariki Act 1989 and the Children’s and Young People’s Well-being Act 1989.
writes of the coproduction between biology and culture that creates embodied experience. She points out that even the most basic of biological events are contingent on the interaction of individual biology with language, cultural understandings and categories of the body, the organisation of societies by class, age, gender, ethnicity, political-economic contexts, and diet or physical environment. Consequently, most experiences of the body—pain, hunger, illness—are variable, though when features of the socio-cultural or physical environment are more or less shared, then individuals may also share aspects of their embodied experience. From these shared experiences of the body, people collectively generate cultural understandings and discourses of health, illness, development, aging and suffering that in turn filter bodily experience. In this perspective, coproduction is constituted through the dialectic interaction of biology and culture which sediment into what Lock terms ‘local biologies’ of bodily experience.

Both authors, therefore, use coproduction to describe the dialectical nature of relationships: for Jasanoff, between science and society, and for Lock, between the body and culture. In this thesis, I employ a similar idea of coproduction to capture the way that children’s health is produced through dialectical relationships between society, the body, and children’s own practices. Importantly, the notion of coproduction offers a way of giving attention to children’s activities as significant and powerful without implying responsibility for their own wellbeing. Victor’s practices contribute to his health: lying to his class allows him to obtain the injections that prevent secondary rheumatic fever recurrence while avoiding the stress of stigma; stealing and sharing food shapes his nutritional status; spitting out his antibiotics alters the microbiology of his body. Yet all of these practices are also structured by Victor’s body, the peer society he co-creates, and wider social organisation of the institution, the community, and the state. He lies to his class in part because of a health system that delivers care through the institution. He shares his biscuits because of a peer culture that demands distribution of resources in a context of scarcity. He spits out his antibiotics because the pills are big for small throats and difficult to swallow, and because the alternative liquid medicine is socially understood to mark a ‘little kid’. Finally, his practices also contribute to meanings within peer cultures—stigma, morality, identity—which maintain, reinforce or transform the social structures that guide his activities as well as those of his peers, teachers, parents, and even policy makers. Coproduction therefore characterises the way that children’s health is made through collaboration between children and society: through practices that are both enacted through bodies and embodied, and structured by society just as they structure future
action. What, then, are the processes through which children coproduce their own health and the health of others around them?

From asthma to rheumatic fever

To answer this question, I spent 13 months, including holidays, driving over an hour to the opposite pole of the city to attend Tūrama School in Papakura, South Auckland. Almost every day, I worked alongside 120 children aged between eight and twelve years (school years 5-8) to research their understandings and experiences of health and illness. As a place where the State directly reaches children’s lives in significant ways, a school setting offered the opportunity to observe how children’s peer cultures interact with government policy to coproduce health.

I was introduced to Tūrama School by a former work colleague, Mrs Randall, who was now a senior teacher there. On my behalf, she asked the principal if I could conduct my research with them while acting as an informal teacher aide, and within a day I had a site for my study. After meeting with the principal and gaining ethics approval, in 2015 I eventually joined Mrs Randall in her classroom where I was based for the first few months of fieldwork, building a solid set of relationships with the 22 children there. After a while, I began to visit six other classes, initially rotating day by day, and later focussing on sustained work with just three, though I still visited the others regularly. I volunteered my services as a ‘classroom helper’ and some teachers took this up more than others. I mounted work, cut up labels, taught art lessons, and accompanied classes on myriad field trips: athletics day, swimming sports day, the museum, the local theme park, end of year trips for the graduating year sixes. I also filled two workbooks with handwriting and mathematics exercises, spelling words and diagrams of volcanoes as I worked alongside children in the classroom, moving around different groups of children as I usually took the spare desk of whomever was away that day.

My original plan was to focus on asthma, a disease which is very common among New Zealand children and disproportionately affects Māori and Pasifika children (Ellison-Loschmann et al. 2009). I proposed to establish myself within a school, observing how asthma plays out within a context of everyday health meanings and practices at school, while also inviting children with asthma and their families to participate in a focussed asthma study.

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6 Though primary school usually ends after year six, children in the bilingual classes may continue up to year eight.
across the year at home. Though I was able to recruit 82 children for the school study, and seven children into the asthma study, it became apparent within the first few months that though asthma was common, it was not particularly salient amongst the complex of intersecting threats to wellbeing and the complicated sets of practices children and parents used to negotiate these. When children are experiencing the demands of poor nutrition, cold, stress, infections, insufficient sleep, bullying, domestic violence, addiction, family illness and social instability, to focus on asthma seemed rather like missing the forest for the trees. Moreover, high rates of rheumatic fever among this group of children meant that Tūrama School, like other low-decile schools with high Māori and Pasifika enrolments, was the target of a government programme which included a public health promotion campaign and a school-based clinic. These interventions were powerful intrusions into school life, and begged attention to children’s perspectives which had until then been neglected in policy.

While I continued working with the families in the asthma study, I initiated a third component to the research, inviting all children, with or without their families as they wished, to participate in an open-ended interview about health and illness more generally, and strep throat/rheumatic fever in particular. An additional 38 children participated, some in pairs or threes, and several, having enjoyed the experience, participated in a second interview. I also interviewed four teachers, the deputy principal, the school social worker, the clinic nurse, and six parents or caregivers, as well as formally or informally meeting an assortment of other whānau [family] members.

How are children actors in their health care?

In writing about New Zealand children, I follow a long legacy of anthropologists researching childhoods and families in the Pacific. This includes significant ethnographic work on child rearing in Aotearoa and Polynesia by Waikato psychologists Jane and James Ritchie, whose publications span almost half a century (examples include Ritchie 1957; Ritchie and Ritchie 1970, 1979, 1997). Throughout this work, the Ritchies argue strongly for a distinct pattern of Māori child-rearing, based on evidence collected over time and space, and are vocal advocates for this pattern to be recognised in policy and social services, and for child and

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7 Although sometimes used by Westerners as a broad equivalent for the English word family, ‘whānau’ and Pacific equivalents designate a much broader group and deeper emotional and tribal allegiance (Ritchie and Ritchie 1979). In modern New Zealand, the inclusive valence of ‘whānau’ has also become commonly used to refer to close-knit communities of people who are not relatives, such as work teams or community groups, indicating a relationship that is ‘like’ a family. ‘Whānau’ can also be used refer to Māori communities more broadly (and can include Pasifika).
Māori rights more generally. Joan Metge also pays attention to Māori children as positioned within the whānau (Metge 1967, 1995). More recently, Relinda Tap writes of Pākehā [New Zealand European] and Dutch childhoods from the perspective of parents (Tap 2007), and a body of anthropological work on children has appeared in relation to health, again mainly from the perspective of parents (Mavoa 2004; Park 2000; Scott, Laing, and Park 2016; Trnka 2017). Across the Pacific, anthropologists have conducted ethnographic work on children and families, particularly debating questions of how children become acculturated adults and how families distribute child-rearing, for example in Samoa (Holmes 1974; Mead 1930), Tonga (Morton 1996) and Fiji (Toren 1990; also Karen Brison’s work on identities (2014)). Given the strong focus on socialisation however, children’s own experiences, perspectives, and cultures have been the subject of less anthropological attention in this part of the world (though Mead (1930) and Morton (1996) both interview adolescents), and to my knowledge, this thesis represents the first ethnographic representation of children’s own cultural production in a Pacific context.

I therefore find my analysis of children’s cultural production and health situates itself within the interpretive childhood literature, where the significance of children’s meaning making to health is well represented by a stream of child-centred studies of illness, largely set in West Europe or North America (Bluebond-Langner 2000, 1978; Christensen 1999; Clark 2003; James 1993; Mayall 1996; Prout 1986, 2000a; Prout and Christensen 1996; but also see: Hunleth 2017 in Zambia). Beginning with Myra Bluebond-Langner’s (1978) ethnography of children with terminal cancer, interest in children’s illness perspectives burgeoned at the end of the 20th century following the emergence of what is sometimes termed the new social studies of childhood (NSSC) and James and Prout’s (1990) call to view childhood as a culturally constructed product of a given society. Central to these studies is the recognition of children’s agency: their capacity to produce their own meanings and practices in relation to illness that are distinct from, but influenced by, adult notions and practices. For example, in an ethnography of children’s chronic illness, Cindy Dell Clark (2003) illustrates the many creative ways that children use ‘imaginal coping’ to transform the discomforts and mundanities of diabetes or asthma into fantasy, play, humour or ritual. Children can also demonstrate their agency by manipulating adult constructs of illness. In a study of a small English community, Alan Prout (1988) showed how children could work with social meanings of illness to influence adults’ decision-making, for example, feigning or disguising symptoms to influence what activities they did or did not participate in.
This movement towards recognising children as social actors has therefore been an important reconfiguring of assumptions about children's role in their health care. However, while numerous analyses consider how children's bodies may affect their experiences and meaning-making, the interpretive literature on child health generally stops short of analysing how children’s practices may impact their biology. Yet asking how children contribute to their bodies reveals the limitations of agency as an analytical concept in the face of another problem: how to position children as social actors without implying responsibility for well-being.

Indeed, a number of scholars have cautioned against using agency too uncritically in analyses of childhood (Ansell 2014; Campbell et al. 2015; Durham 2008; Lancy 2012; Mizen and Ofosu-Kusi 2013). Interpretive studies of childhood can tend towards a romanticised depiction of resilient and resourceful children making creative choices to overcome adversity, while masking the constraints on their choices, the extent of their suffering, and the conditions under which children do not have agency (Prout 2000a). Campbell and colleagues argue against this tendency to view agency as a child's ability to engage in any form of action per se, referencing Andersen's (2012) point that agency is 'a blunt analytical tool to describe a 12-year-old girl's choice to have unprotected sex with an HIV-positive older man, to generate income to feed her younger siblings' (c.f. Campbell et al. 2015:55).

Yet studies of how children’s health is produced have tended to avoid the issue of how they use agency, pointing upstream to the historical, social, and political-economic factors and cultural ideologies which structure people’s lives and practices. The power of global political, economic and social forces to shape childhoods has been well recognised in anthropology (Scheper-Hughes 1996; Stephens 1995). Critical studies of childhood, like Kristen Cheney’s (2017) analysis of children orphaned by the HIV/AIDS epidemic in Uganda, consider the impact of inequality, from local to global scales. A body of research in biological anthropology considers the impact of policies and structures on children’s bodies, where biological outcomes such as physical growth or morbidity rates are employed to measure the effects of disparities in exposure to risk or access to care and material resources (Bogin and Loucky 1997; Panter-Brick 1998b). However, in this literature children tend to be positioned as passive experiencers and absorbers of the conditions that impact them. In his chapter on childhoods in the crackhouses of East Harlem, for example, Philippe Bourgois powerfully describes the children he encounters as becoming ‘ground up’ into the underclass (1995:261) as they internalise the conditions of El Barrio. Yet as those interpretive studies of childhood
agency have demonstrated so emphatically, children are also engaged in the production of culture, so the concept of internalisation is not sufficient to explain the processes of social reproduction. How, then, to position children as contributors to health when much of their health status is, like Victor’s, the product of powerfully converging historical, social, and political economic forces?

To resolve this, I draw on the theoretical work of sociologists William Corsaro (1992, 2015), Pierre Bourdieu (1977, 1984), Chris Shilling (1993) and Alan Prout (2000b, 2005) to unravel the way children’s practices contribute to the coproduction of their health through their engagement with and transformation of the social structures that guide and constrain their practices. These scholars all deal with issues of structure, agency, and the body in social reproduction. While Bourdieu is more focussed on the structures and outcomes of socialisation in the form of the internalised habitus, Corsaro does the work of isolating the mechanisms of socialisation and expanding these beyond internalisation to children’s cultural appropriation, reinvention and reproduction. However, Bourdieu’s conception of embodied practices constructing and constructed by social structures brings a necessary view of the way children’s interpretive practices are constrained and guided by the circumstances within which they are produced. Shilling, also drawing from Bourdieu, elaborates on how these practices articulate with biological bodies, and Prout extends Shilling’s biosocial theory to children, laying the foundations for understanding the dynamics of children’s participation in the coproduction of their health. Though they may not use the term ‘coproduction’, all of these scholars are concerned with how individuals—or individual bodies—and society produce each other, and thus their insights contribute to my analysis of how children, together with society, come to coproduce their health.

Each of these theorists emerge from concurrent, distinct, though interconnected lineages of 20th century scholarship across anthropology, sociology and psychology. In the remainder of this chapter I therefore trace the ways that questions of children and social reproduction have been dealt with through parallel lines of enquiry through socialisation, practice theory, and developmental psychology. In particular, I note four major shifts in thinking about childhood which lay the groundwork for this thesis: first, the movement towards recognising children’s agency within their socialisation; second, the understanding of agency as working dialectically with structure; third, the extension of socialisation processes beyond individual internalisation to a collective and transformative reproduction; and finally a view of the body as biosocially produced.
Finding agency in socialisation

The term socialisation is used to describe the processes through which individuals—particularly children—become members of society and society reproduces itself. Interest in these processes developed across the social and psychological sciences of the early 20th century United States, where the decline of child mortality due to public health measures and mass enrolment in secondary schooling generated new attention to childhood through what LeVine calls ‘pediatric and pedagogical terms’ (2007:248). In psychology, this meant the emergence of modern developmental psychology, which sought to understand the cognitive and socioemotional processes through which children grew into adults. For anthropology, the subsequent wave of psychological theory around the child, including the theories of Freud, Piaget, Vygotsky, and Bowlby, generated much of the agenda in the United States, and a proliferation of child-rearing and cultural transmission studies documented the variation in socialisation practices and effects across cultures (Montgomery 2009). These studies included the influential Culture and Personality Studies led by pioneering anthropologists such as Ruth Benedict, Margaret Mead, and Edward Sapir, which set out to address empirical questions about the relationship between childhood experiences and enculturated adults (Schwartzman 2001). In general, these studies and those that have followed are concerned with the ways in which adults mould children into particular kinds of social citizens; the child here is mostly passive, but embedded in social relations and interactions.

To this day, anthropologists remain interested in the processes of socialisation (Bolin 2006; Briggs 1970, 1979; Broch 1990; Geurts 2003; Morton 1996). In a recent example, Bambi Chapin (2014) makes a return to culture and personality, using psychoanalytic approaches to investigate questions about the processes of socialisation in Sri Lanka, via relationships and under the influence of religion, which lead to a cultural deference to authority from children as young as 10. However, the literature has evolved, in part under the influence of the new childhood studies in the early 1990s which emphasised children’s agency, with an expansion of socialising agents from the parent to the wider family, including children themselves, the community and peer group. For example, Brigg’s (1998) study of Inuit socialisation follows a single three-year-old child, positioning her as an active agent in every interaction documented, and inferring her perspectives where possible. Such a reconsideration of children’s agency in socialisation processes also emerged from the language socialisation subfield of linguistics, which criticised traditional anthropological socialisation for treating children as the passive recipients of culture and overlooking how everyday language
functions as a key medium of socialisation. Instead, scholars working from a language
socialisation paradigm examine how the processes of linguistic and cultural development are
interlinked, and highlight the active role of the child in acquiring and generating language
(Heath 1983; Ochs 1993; Schieffelin 1990; Schieffelin and Ochs 1986). Meanwhile, some
branches of childhood anthropology, such as studies of cognition and learning, have moved
away from socialisation frameworks, instead following the developmental psychology of
Piaget and Vygotsky to centre the child as the agent acquiring the understanding. For
example, Christina Toren (1993, 1999) explicitly rejects socialisation as something adults do
to children, instead arguing that although others help create the structures, it is the individual
who constitutes their own meanings from these, what she calls human autopoiesis, or self-
making.

More recently, Allison James presents an extensive ‘child-centred’ study of socialisation
from children’s own perspectives using Smart’s (2007) frame of the personal life (James
2013). Asking what socialisation from a child’s perspective would look like, James bridges
the large gap between the decades long concern with child rearing for social reproduction and
the NSSC’s preoccupation with agency. She explores how children experience, perceive,
negotiate and transform various traditional institutions of socialisation—the family, the
school—in collective and embodied ways. This creates a shift from seeing the family and the
institution as socialising children, to seeing how, through their involvement in the practices of
these institutes, children become socialised. As James engages in detail with practice theory,
this work also represents a shift from the original childhood studies to view when, and how,
children’s agency is subject to constraint. This is a version of children’s agency that is
contingent and negotiated, working dialectically with the structures of the family, the
institution, and society, coproducing childhood.

Theorising socialisation and social reproduction

While the anthropologists of the mid-20th century tested psychological theories with
ethnographic accounts of childrearing across cultures, within this literary milieu sociologists
also developed influential theories of socialisation and social reproduction. Perhaps most
significant of the early theorists is Talcott Parsons (1951), who drew on Durkheim to answer
the Hobbesian problem of how order is maintained in society. For Parsons, the notion that
social systems were entirely maintained through rules and laws was insufficient, as he noted
that individuals tend to work in ways that sustain, rather than destabilise systems (James 2013). Society does not simply impose social rules, Parsons argues, instead, families and schools play a functionalist role in transmitting culture and practices to the next generation, whereby from a young age, children begin to internalise social norms to the point where they become self-imposed.

Among other issues, it is this uninterrogated ‘internalisation’ concept that Dennis Wrong particularly critiques in what he labels an ‘oversocialised view of man’ (1961:184). Wrong argues that Parsons has misconstrued the original Freudian meaning of the ‘internalisation’ term to loosely mean ‘learning’ or ‘habits’, with the implication that the individual affirms as well as conforms to those norms. This neglects the ‘inner-conflict’ aspect of Freud’s superego, which suggests only that an individual will feel guilty for not living up to a norm—not that they actually will conform. Internalisation, as it is used by Parsons, is therefore a taken-for-granted assumption of fact that, without an unpacking of the underlying mechanics, has limited explanatory power. In fact, this is just a reversal of the question, Wrong points out, writing, ‘How is it that violence, conflict, revolution, and the individual's sense of coercion by society manage to exist at all, if this view is correct?’ (1961:186). What are the circumstances under which such ‘internalisation’ does not, or does not fully, occur?

Such opposition to the Parsonian view of individuals internalising the rules of society, as well as the constraint-based theory of scholars concerned with hegemonies of power (Karl Marx, for instance), cognitive structures (Claude Levi-Strauss) or cultural systems (Clifford Geertz), gave rise to the practice theory of the 1970s and 80s and its attempts to explain the relationships between human action and ‘the system’ (Crossley 2001; Ortner 1984). Practice theorists such as Bourdieu (1977), Giddens (1979) and Sahlins (1981) argued for a dialectical, rather than oppositional relationship between agentive practice and constraint; while social structures and systems constrain actors, they are also produced, reproduced, and can be transformed through practice—what people do (Ortner 2006).

I focus here on Bourdieu in no small part because the embodied nature of habitus has important implications for the present study of health. Bourdieu’s theory of practice involves individuals as social actors deploying different forms of ‘capital’ (economic, social or symbolic assets) in strategic negotiations across different ‘fields’, or distinct domains of action such as home, work or school (Bourdieu 1977, 1986). Central is Bourdieu’s concept of habitus, an enduring and transferable set of schemes which are internalised within individuals through the repetitious practice of certain modes of being, thinking, and behaving which
sediment into largely unconscious, predisposing structures for future action (Wacquant 2005; Maton 2008). In other words, people become what they do and do what they become. What people do, however, is also guided by pre-existing historical, social and environmental structures such as class and material opportunities which constrain, or constrain perceptions of possible action. For example, Connolly (2004) describes how children who succeed in school tend to come from middle class homes where they have acquired the ‘dispositions’ and educational capital from parents who themselves value and encourage engagement in education and possess the forms of knowledge hegemonic in the national curriculum. These processes explain the tendencies of gender or class structures to reproduce themselves, as individuals experiencing common conditions (and conditionings) will share characteristics of the habitus but also perceive these commonalities to reflect the natural or commonsense order of things (doxa) (Bourdieu 1977). Yet while Bourdieu’s habitus has, at times, been criticised for being overly deterministic in that individuals are born into a structured world that predates them (Jenkins 1982), Bourdieu conceptualises these matrices as circular and evolving; the structured world has been constructed by prior individuals, and will form a base from which future individuals will reproduce, innovate, and generate new structures that predispose, rather than predetermine future action (Crossley 2001).

Childhood is a period of critical importance for Bourdieu as within early childhood the primary habitus is laid down, setting the trajectory for (though not wholly determining) an individual’s lifelong embodied practice (Bourdieu 1984). In recognising the ‘structuring activity of agents’ (1986:467) Bourdieu leaves open the possibility that children too could be agents who engage in creative praxes that generate and modify habits. However, Bourdieu and colleagues pay little empirical attention to children or the precise processes of that early cultural practice, instead largely outsourcing those explanations back to socialisation and the internalisation concept (Garnier 2015), as well as pointing to formal education as a strategy through which families can reproduce their social position in subsequent generations (Thomson 2008). Although documenting in detail the processes of remodelling a secondary habitus through a boxing gym (Wacquant 2004), for example, Wacquant describes the process of acquiring the primary habitus vaguely as occurring ‘through osmosis’ in the family microcosm (2016:68). Thus Bourdieu and his contemporaries leave open a theoretical space for childhood theorists to grapple with.

Indeed, because of childhood’s relevance to Bourdieu’s theory of habitus, a number of childhood and education scholars have elaborated Bourdieuan theory to understand aspects
of childhood and schooling (Alanen, Brooker, and Mayall 2015; Connolly 2004). A large emphasis has been on reconstituting Bourdieu’s (1977) concern with patriarchal relations of power within intergenerational analyses of social change and dominance hegemonies that structure and maintain inequalities between adults and children. For example, Vuorisalo and Alanen (2015) analyse how preschool children use strategies to acquire and deploy different forms of capital in negotiation with teachers, and how this unequal distribution of capital results in an unrecognised, stratified social order—a form of symbolic violence. A second prominent theme has been the effects produced by incongruities between the forms of habitus generated in different fields, whether it is the impacts of differences between home and school habitus on educational achievement (Connolly 2004) or parent-child negotiations (Mayall 2015), or how conflicts between local and globalising ideas about childhood shape children’s dispositions for work and labour practices (André and Hilgers 2015).

I extend these ideas in this thesis, considering how new kinds of health services, ideologies, changing demographics and patterns of morbidity and mortality can contribute to new kinds of struggles in the field and through new practices restructure forms of habitus for the children who negotiate these changes in school contexts. The emphasis in childhood scholarship, though, tends to be on how children negotiate interpersonal relations, or how changing social and political-economic environments engender generational differences in dispositions. Less attention is paid to the processes through which children collectively produce cultural meanings and practices, the ways in which they deploy cultural forms to navigate intergenerational, institutional, and political-economic constraints, and how the embodiment of such practices helps to reproduce inequalities. In this thesis, I use a view of children’s socially structured cultural production to reveal, for example, how children establish and embody practices of ‘not hungry’ rather than accept stigmatised provisions, or how children construct an understanding of rheumatic fever aetiology which shapes their perceptions of risk and subsequent practices of accessing health care. This understanding of children’s collective cultural production comes out of a different way of conceptualising childhood learning; the shift in thinking brought by the developmental psychologists who laid the groundwork for Corsaro’s (1992) model of interpretive reproduction.
Developmental psychology and interpretive reproduction

The constructivist approach of developmental psychology challenged the determinism of early socialisation theory, instead emphasising the child’s active role in socialisation processes. If, as Corsaro (2015) puts it, for the social determinists society appropriates the child, then for the constructivist developmental psychology it is the child who appropriates society, albeit still primarily through processes of internalisation.

Of great significance from this era is the work of psychologists Jean Piaget (1967) and Lev Vygotsky (1978), the latter of whom directly influenced Corsaro’s (1992) model of interpretive reproduction. Piaget has been the more well-known, although he began writing in the 1920s and it took several decades for his work to gain influence. By the 1960s, Piaget’s theories dominated developmental psychology. His legacy is the view of children as active participants in their development, and the argument that from infancy, children interpret, organise and use information from their environment to construct conceptions (mental structures or schema) of their physical and social worlds (Ginsburg and Opper 1988). This is a process that primarily occurs inside individual children’s heads, constrained by the biological development of cognitive structures. Cognitive development in Piaget’s view therefore occurs from the ‘inside out’ (Burman, c.f. Connolly 2004), first in the individual minds and then expressed ‘outside’ through activity and behaviour.

By contrast, Vygotsky (1978) situates children’s development within their collective actions and interactions set within society. Development, in this view, instead occurs from the ‘outside in’, whereby a child first experiences the world as an actor embedded within culture and social relations, and these interactions secondly become internalised as mental schemes of social and physical systems (Connolly 2004). Language is therefore a critical part of children’s development, first used to communicate with others, and then as internal speech or thought used for self-regulation and consciousness (Smith 2013). The significance of this process for learning is captured in the ‘zone of proximal development’ concept, whereby through social interaction children can engage in more complex forms of activity and behaviour than what they could understand or accomplish on their own (Vygotsky 1978). While Piaget saw children’s capacities as limited by the biological architecture of brain development, for Vygotsky the limitations can be found in what, when and with whom children experience, or what more recent scholars have termed the quality and quantity of
‘scaffolding’ others make available to the child to support them in stretching their capacities (Smith 2013).

Although Vygotsky died in 1934, his work only became influential in Western scholarship from the 1960s and 1970s, and in recent decades has been particularly taken up in education fields. For example, Paul Connolly (2004), who tackles the increasingly global question of boys’ underachievement at school, finds value in Vygotsky’s attention to the social nature of learning, which has particular relevance for children’s developing ideas of gender and class. However, Connolly develops Vygotsky’s work further by considering how children’s socially learnt ideas and behaviour are influenced and shaped by cultural contexts and differential relations of power embedded in much broader communities, incorporating insights from Bourdieu and Elias. While scholars in the Vygotskyan tradition typically focus on simplified two-way interactions between a child and one other individual, using Elias’ (1978) ‘figuration’ concept Connolly locates children within open, interdependent networks of relations that, crucially, are infused with differential relations of power that create systems of dominance and determine which ideas are established as norms. Secondly, Connolly finds common ground between Vygotsky and Bourdieu in the processes of internalisation; the way that cognitive schemes, once internalised, form the habitus which structures future action and reflects the particular contexts and broader social structures of children’s lived experience.

The work of both Vygotsky and Connolly, however, still remains uncritical of the reliance on internalisation as the key mechanism of cultural production and learning, which limits children’s activities to the acquisition of adult culture and practices, measuring child competencies and understandings against an adult standard and emphasising outcomes over processes. Corsaro, while heavily influenced by Vygotsky’s notions of children’s development as collective and interactional, constructed his (1992) model of interpretive reproduction by extending processes of social reproduction beyond simply internalisation to include children’s appropriation, innovation, and creative interpretation of culture. Like Vygotsky, Corsaro breaks away from theories of socialisation which position the child as individually and privately internalising the skills, knowledge and culture of adults in a linear fashion. Instead, he views childhood socialisation as a collective, social process where children remake culture anew.

Using the metaphor of an orb spider web, Corsaro (2015) describes how children work from the family unit, represented at the centre of the web, and collectively spin spirals of peer cultures across the spokes of the web, which represent different fields or locales that compose
various social institutions (such as family, educational, religious etc.) Through these peer cultures, knowledge and practices are produced, reproduced and gradually develop into forms that enable participation in the adult world. Focussing on play within children’s peer group, Corsaro describes how children appropriate adult cultural frameworks and routines and transform these into variations, for example through ‘keying’, where a new activity is modelled on the patterning of an existing activity, or through ‘embellishments’ where certain elements of routines are intensified through repetition or exaggeration.

Interpretive reproduction offers a way of viewing children as significant contributors to health through their collective production of cultural meanings and practices, for example, the way that children construct novel concepts of sore throat, or the ways that adult frames of death influence, but are not duplicated in the way children themselves employ death concepts and language. While Corsaro’s model is strong on agency, it is, as James (2013) notes, weak on constraint. Structure is present, represented in the spokes of the orb web around which children spin spirals of meaning. But children’s activities are not a free-for-all of endless possibility and the model does not capture the particularities of how these structures guide children’s cultural production: how options are constrained by relations of authority or resource availability; how the body itself structures cultural production as the material substrate through which sociality is enacted and embodied; and to paraphrase Clifford Geertz (1973), how children become caught in the webs of meaning they themselves have spun.

In this thesis I therefore aim to work in the gap between Corsaro and Bourdieu, detailing the way children’s interpretive practices articulate with both the biological structures of a body and the constraints of material deprivation and institutional and societal structures. I argue that these processes are best captured by a coproduction framework which makes room for children’s agency by positioning them in dialectical relationship with the body and society: each reinforcing, remodelling, or transforming the other. Conceptualising children’s activities as a coproduction shifts interpretive reproduction from webs made from children’s spirals creatively freewheeling over adult spokes, to seeing children’s cultural meanings and practices as constrained and enabled by experiences of the body and wider social systems, while those meanings and practices simultaneously mediate how those wider social systems work on bodies. Coproduction also means that children, in negotiating the structures of their childhoods, collectively create new cultural forms within peer groups that not only help to structure their own habitus, but leave open the possibility for children to modify the habitus of their parents as well.
Why is the body important?

In many studies of biological or medical anthropology, children’s bodies have represented important physical records of environments and events experienced, encoded in measurable processes of growth, development and repair, biochemistry and disease (Flinn 1999; Horton and Barker 2010; Littleton 2007; Panter-Brick 1998a; Panter-Brick, Todd, and Baker 1996). The role of the body in these studies is to provide a map for the distribution and dynamics of power and resources within and across populations (Krieger 2005). However, the body is not only the material substrate upon which structural inequities leave their marks, but the locus of experience (Merleau-Ponty 1962) and of practice (Bourdieu 1984)—the sites at which the structured world is converted into social meanings and translated into practices. Children’s interpretations are important mediators of their own health because their practices have real physical effects. For humans, the embodiment of inequity, therefore, does not only involve the biological processes of converting energy to growth, immune function and homeostasis, but experiences of the body incarnate (Frankenberg 1990) made meaningful in different ways within unequal social worlds.

The body already underpins childhood, which is socially constructed around a universal period of biological immaturity, the significance of which is culturally variable (James and Prout 1990; Prout 2000b). In the West, this immaturity is conflated with social vulnerability and incompetence, producing childhoods characterised by a prolonged period of dependence and restriction. In other places, such as southeastern Mexico (Kramer 2005), and Java and Nepal (Nag et al. 1978), incomplete growth does not preclude the expectation of being a full and contributing member of society, and children’s labour activities can form an essential part of domestic and community economic life. Within this period of growth also sits a number of reasonably canalised biological life events upon which variable degrees of significance may be socially ascribed by adults and children. The eruption and loss of teeth are socially celebrated by the tooth fairy in North America (Clark 1998) or mark a new life stage for the Ngoni children of Malawi (Read 1968). While Davis and Davis (1989) suggest there was virtually no adolescence in traditional rural Morocco as onset of puberty is quickly followed by marriage and the conferring of adulthood, in Aotearoa and elsewhere in the West, neurological science is used to extend the stage of perceived immaturity up to 25, with a host of subsequent policies and interventions (France 2012).
The body, therefore, bears important social markers which differentiate and transform social experiences across societies, just as these experiences in turn transform bodies. These markers of growth, however, along with other universal, but culturally variegated bodily experiences—illness, injury, fatigue, emotion, stress, hunger—are biosocially produced; the underlying biology itself is underwritten by cultural processes which mediate how these experiences are distributed and interpreted. Thus, sociologist Chris Shilling (1993) describes the relationship between the body and culture as dialectical—a coproduction where each simultaneously helps to produce the other.

Shilling’s ideas developed from Turner’s (1984, 1992) concern that sociological treatments of the body have tended to fall into either biological or cultural reductionism, what he terms ‘foundationalist’ and ‘anti-foundationalist’ approaches. Foundationalist approaches assume the body as a material object, distinct from the social, while anti-foundationalist approaches claim that it is not possible to make a distinction between the body and its representations, since we only have access to the materiality of the body through the discourses which structure and shape it. Turner proposes a reconciliation of these two approaches by way of ‘methodological eclecticism’ (Prout 2000a:4); acknowledging the value and intellectual validity of both approaches, and seeing them as complementary. Shilling, however, argues that combining the two approaches ‘without altering any of their basic parameters’ (1993:103) is not theoretically coherent, and that it is the relationships between the body and society that are critical. He therefore proposes a framework which takes the body as a social and biological work in progress throughout the life course, with each resourcing and constraining the other; a dialectic coproduction. For example, Shilling describes the embodiment of social gender inequalities which occurs where average biological sex differences are socially highlighted and transformed into absolute and naturalised gender differences. The subsequent socialisation towards particular patterns of behaviour further reinforces those differences: encouraging men to build muscle mass, and women to maintain small bodies. The biological and the cultural thus work together to mutually coproduce both the body and society.

Despite the importance of childhood as the foundational stage of this biosocial development, Shilling’s (1993) discussion of childhood is limited to chapters on Elias and Bourdieu, where, as noted in the previous chapter, children are positioned as passive receptacles of

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8 Bourdieu similarly describes an embodiment of gender in terms of tastes and bodily expressions (1984:190–92)
socialisation and ‘civilising’. Instead, sociologist Alan Prout builds on Shilling’s thinking to theorise the dynamic relationship between the body and culture in childhood and challenge the biosocial dualism that had separated the social constructivists from the developmental scientists (Prout 2000b, 2005). In this view, the body is the site of children’s experience and their environments both structure and are structured around their body; children are therefore hybrids of their biological inheritance and the social world which together mediate their cultural and biological development. This is an understanding that has long been present in anthropology: Boas’ early recognition of the biocultural nature of developmental plasticity set the stage for a proliferation of cultural studies of childhood (Boas 1974[1911] c.f. Montgomery 2009), where the moulding and disciplining of children’s bodies in culturally patterned ways is a main feature of socialisation (e.g. Geurts 2003), while child-rearing practices are configured around the biology of infant bodies (LeVine 1977). This attention to the body carried through to the interpretive childhood anthropology of the early 1990s which further located the body within children’s experiences and understandings of identity and illness (e.g. Christensen 1999; James 1993). However, Prout’s notion that children (and therefore adults) develop within a dialectical ‘medley of culture and nature’ opens up a space to conceptualise the role children play in the production of their own bodies—bodies not only as made by adults and experienced by children, but as a coproduction between children and society, biology and culture, agency and constraint.

Prout’s work has sparked new theoretical interest in the hybridity concept: to break down nature/culture binaries in childhood studies (Kraftl 2013; Ryan 2012), navigate the biopolitics of childhood (Lee and Motzkau 2011), reassert children as biological creatures within a changing environment that has particular effects upon those biologies (Lee 2013), and open up childhood studies to more cross-pollination with the psycho-biological disciplines (Thorne 2007). Very little of this recent work has been ethnographic, although Davies (2015) emphasises the embodied sensory and inter-physical nature of children’s family relationships in a context of socio-economic deprivation in central England, showing for example, how children monitor the bodies of younger siblings for care needs, and negotiate power hierarchies through physical confrontation. Still missing, however, are ethnographic accounts of how children coproduce their bodies in relationship with society, and how these biosocial processes of the body and its interpretation unfold, temporally and in context.

What then, are these processes through which children co-construct, reproduce, internalise, and embody their worlds? How are children’s health practices structured by institutions, state
interventions, wider cultural discourses and material resources? How does material deprivation become translated into interpretive practices of the body; a habitus of childhood inequality? How do children negotiate, accommodate and embody structural vulnerability produced by their social positioning? And how does the instability and risk brought by structural violence become woven into children’s cultural production?

These are the questions I will address through the remainder of this thesis. They are questions less about what children experience, and more about what children do—though, of course, children’s practices derive from their experiences. Importantly, these questions point to why what children do matters—they suggest that the child who is invisible in policy is very much active in the coproduction of their health. These perspectives reveal children who are actively negotiating unstable, constrained worlds, but worlds that are in part produced by children themselves, as they collectively translate their material conditions and cohabitants into cultural meanings, social relationships and embodied practices, at the same time as they are shaped and constrained by those conditions. As economic inequalities stamp biology with signatures of poverty—respiratory illness, dental decay, stress—the state targets these young bodies and through their experiences of health interventions in the institution, children construct new understandings of illness and generate new practices of the body. This is a view of how children co-create their habitus, working creatively within structures which open up or constrain opportunities and resources, and in doing so help to generate the habitus that in turn guides future practice.

Overview of chapters

I address these questions through the following chapters. In chapter two I establish the world of Tūrama School, both through broader political-economic, historical and societal perspectives, and as seen through children’s own eyes. This institutional environment, shaped by local, national and global forces, set the structures that children negotiate and transform in the coproduction of their health.

Chapter three sets out the main power structure operating within the school, the separation of adult staff and child pupils, which explicitly works on children’s bodies differently from adults. The dynamics at work here were keenly felt by myself as a researcher, whose adult body transgresses these boundaries and inadvertently reveals the power of these forces. In this chapter I discuss how such institutional structures challenge the role of the researcher
who attempts to balance what are sometimes competing ethical obligations to children and adults, and propose the role of ‘transparent guest’ as a guide for navigating between adult and child worlds and their associated rules and expectations.

Having established the geographic, social, historical and political economic contexts for this thesis and the methodological complexities within which the data were produced, over the following five chapters I unpack the dynamics of health coproduction for children at Tūrama School to answer the question: how do children participate in the coproduction of their health? I take the case of the school rheumatic fever prevention clinic and related campaign as my point of entry in chapter four, showing how children collectively create cultural meanings of illness and health care routines around the clinic in ways unanticipated by policy-makers. Children’s embodied interpretations of illness and creative practices mediate their engagement with health care and pharmaceuticals, key ways that they shape their health in coproduction with health care services.

In chapter five I consider how children coproduce their health together with state discourses of responsibility. Focussing on public debates around the provisioning of school lunches, I describe how children contribute to their dietary patterns by making social meanings and practices around different kinds of food, shaped by wider discourses and material constraint. In chapter six, this material deprivation becomes translated into interpretive practices of the body to form a habitus of childhood inequality which in turn orients future practice. I argue that structural constraints on resources or power shape the degree to which children tune in to particular bodily signals which tell children they are cold, hungry, or sick. At the same time, children’s peer cultures produce socially accepted meanings and practices which further reinforce how those signals are experienced in the body.

In chapter seven I explore how the instability wrought by economic and housing policy is translated into children’s practices of resilience, and how the interconnected and embodied nature of these practices contributes to children’s health. I draw on socio-ecological frameworks for resilience which reconcile children’s structural vulnerability with their competence as social actors by placing the child within broader social and physical environments which constrain their activities or provide resources for children to use. Finally, in chapter eight I consider how children coproduce their health in relation to the structural violence which mediates their experiences of death. Tūrama School children’s ‘small talk of death’ can reveal their experiences of a social positioning where they are thrice marginalised by their ethnic, socio-economic and age statuses, the confluence of which creates a context
where death is both common and culturally salient. While children collectively compile understandings of what death means and the likelihood that it will affect them, adults invoking death, for example in health promotion, inadvertently tap into a powerful expressive device of children, both reinforcing a sense of fear and urgency around death, and shaping children’s practices of health care.

Together, these chapters illustrate how the coproduction of children’s health occurs within peer ecologies, grounded in the body, and guided by wider institutional and societal structures which children translate into their own collective meanings and practices. I conclude in chapter nine by reflecting on the implications for theory and practice of conceptualising child health in this way: as a coproduction between interconnected, embodied individuals, making meanings from their experiences of a structured world, collectively practicing childhood.
In one of my first conversations with Ruby, she asks me if I want to be in a gang.

‘Uh, no,’ I say, emphatically.

‘I want to be in a gang when I grow up,’ Ruby tells me. Her father is in a gang. ‘I want to be like my Dad. I don’t want to be like my Mum, she does nuffink all day.’

‘Is there anyone else you could be like?’ I ask hopefully, and Ruby pauses, thinking.

‘Mm, there’s my auntie. She works at Countdown!’ she says dubiously. ‘But I think being in a gang would be better.’

While teachers described Ruby’s gang-member father as a scary man, I could see why they tended to have a soft spot for Ruby. Mrs Randall, who taught her in year five (age nine), told me she initially thought Ruby was slow, she was so withdrawn. By year six, the green-eyed, ruddy-cheeked Ruby had elbowed her way into position as the undisputed leader of the girls in her class. ‘She’s the boss of us. She’s like our mum,’ her friend Marielle tells me one day, explaining how she and the other girls have to ask Ruby if they want to go to the toilet, because she wants to know where they are going ‘or else we might get hurt’. As a ten-year-old, Ruby is bold, insightful, and, though I once spotted her changing her maths score from a 98 to a 100, academically confident.

I do not believe Ruby at first when she tells me she lives with 18 other people, because she can be ambiguous with the truth at times; she made up a ‘fake’ birthday in December because, she later rationalised, she shares her June birthday with an uncle and wanted her own day. But she draws me a map of the six bedrooms where her grandparents, parents, siblings, aunts and cousins live, plus the sitting room where two uncles sleep. Another auntie stayed the other night, too, because the police were down the road looking for synthetics [drugs], and her auntie was scared—she had been to jail before. To be sure, I later checked the story with her teacher Mrs Charles, who said ‘that sounds about right.’ Mrs Charles

1 The local supermarket chain.
knows a lot about her pupil’s home lives, and she has given Ruby her phone number in case she needs it one day.

Ruby’s circumstances, while not those of an average Tūrama School child, are unusual more for the size of the rental home rather than the people-to-bedroom ratio. Her friend Mila similarly shares a bedroom with three brothers. Ruby’s childhood is, however, far from the average New Zealand childhood. Data from the 2013 census shows the average household size is 3.5 people in Ruby’s suburb, well above the national average of 2.7 (Statistics New Zealand). The crowded conditions likely have various impacts on the family’s health, not least of which is stress; Ruby describes living with so many people as ‘full’ and ‘annoying, ‘cause there’s a lot of arguing’. In an English context, Hayley Davies (2015) found that her child participants experienced overcrowded homes as stressful. Also in this house, Ruby’s young cousin developed rheumatic fever, which is linked to overcrowding (Jaine, Baker, and Venugopal 2011), and he now receives the same monthly prophylactic penicillin injections as Victor, who we met in chapter one.

Tūrama School represents children at the most deprived end of the socio-economic scale, childhoods geographically bounded to the point where, until recent media attention, the deep inequalities differentiating New Zealand childhoods were near invisible to the middle-class. These deprived childhoods also disproportionately belong to Māori and Pasifika children (Henare, Puckey, and Nicholson 2011; Perry 2016). Aotearoa, is a multi-ethnic nation located in the Pacific and administrated by dominant British colonial systems of governance in partnership with the indigenous Māori peoples. Since World War Two, high numbers of Polynesian migrants have come to settle particularly in the South Auckland region alongside urban Māori, and Auckland is now often referred to as the Polynesian capital of the world (Barbera 2011). In Aotearoa, these Māori and Pasifika childhoods are configured around Western legal, political-economic, institutional and social frameworks of childhood which see non-Pākehā\(^2\) children as deviations from the norm, historically represented as novelties for tourists or as mortality statistics, as problems to be ‘Europeanised’ or who need ‘catching up’ to Pākehā level (May, 2001). Māori childhoods are colonised childhoods, and a misalignment between indigenous and Pākehā conceptions of childhood and practices of

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\(^2\) Pākehā is a Māori word which has entered mainstream New Zealand English and generally refers to a New Zealander of European ancestry. Though widespread, the term is contested; it featured in the 1996 census alongside New Zealand European, the official census category, but was subsequently removed, to ongoing controversy as many white New Zealanders prefer to identify as Pākehā. Use of the term is a relatively recent development, which King (1985) argues emerged in response to the Māori renaissance of the 1970s and 1980s, and implies European ancestry but indexes relationship to country as one of the ‘founding peoples’ of the new nation state and relationship to Māori as the other.
child-rearing, embedded in law, policy and institutions, and compounded by neoliberal economic and housing policies has perpetuated inequities and cultural alienation for children like those at Tūrama School. These are the other, the othered Aotearoa childhoods: bicultural childhoods, childhoods in poverty.

From the perspective of Tūrama School children living them, however, these are normal childhoods filled with fun, freedom, and lots of love, even if sometimes things are not so great, like when Ruby worries that her Dad will get shot, or when Whetu gets taken away from his mum, or when Arya gets teased for being poor, even though she is definitely not. The census data that demonstrates the ethnic composition and relative poverty of Tūrama School children is only one representation of a lifeworld that is understood quite differently by those living within it.

Yet economic insecurity, neoliberal reform, cultural dissonance and the legacies of colonisation and migration histories have all contributed to the conditions of Tūrama School childhoods, even if these remain outside the children’s field of view. Historical and contemporary housing, economic, health and social policy shapes both the marginalised circumstances of their families as well as the role of the institution in mediating the harmful impacts. The discourses, practices and interventions of families and the institution in response to economic insecurity comprise the structures that Tūrama School children negotiate, reconstruct, and transform in the coproduction of their health.

In this chapter, I aim to set the scene for the analysis of children’s health understandings and practices to follow by outlining the major structural features of Tūrama School childhoods that form the context for their health coproduction. As the central question of this thesis requires a view of how children come to understand health through their own embodied perspectives, I represent Tūrama School childhoods as they are viewed from historical, political and societal perspectives—the researcher’s view—but also how Tūrama School children see themselves. I consider the geographic and ethnic composition of Papakura, where Tūrama School is located, and how this translates into children’s perceptions of their world, and the impacts of economic inequities that leave 100,000 Aotearoa children in poverty. These inequities are shaped by colonisation and migration histories, and so have significant implications for Māori and Pasifika health; in particular rheumatic fever represents an important contemporary manifestation of eroding housing and economic conditions in South Auckland that can be traced to historical circumstances. Finally, I outline the historical and present roles of the New Zealand state and school institutions in caring for
children’s health and welfare; roles which directly shape Tūrama School children’s health experiences in the classroom.

What culture are you?

The children of Tūrama School have been relegated to an area of highest deprivation within Papakura, South Auckland, through the doubly-marginalising intersection of social class and ethnicity—the result of historical colonisation and migration events compounded by contemporary neoliberal policy. Formerly recognised as its own city and later own district, since the amalgamation of cities into the Auckland super city, Papakura is now officially designated as a Local Board area, with a population of around 45,000. Locals call themselves a town. In this town there is literally a wrong side of the tracks, a train line dividing low-quality state housing from new developments which attract middle-class commuters as a housing crisis drives up prices in the central city. This class segregation is also highly correlated with ethnicity. Māori represented 28% of the Papakura population in the 2013 census—the third highest Māori population in Auckland—and Pacific peoples represent about 14.5%. However local school rolls show a further clustering of ethnic populations; over 70% of children at Tūrama School are Māori (though many of this group will have Pasifika and/or European heritage as well), and around 20% are of Pacific Island descent. 3 Although Pākehā, or New Zealand European people are the majority ethnic group in Papakura as well as nationally, less than 4% of Tūrama School children identify as Pākehā. Meanwhile, schools in more affluent areas of Papakura tend to have few children of Māori or Pacific descent—less than 10%—and higher numbers of Asian and Indian immigrants as well as a Pākehā majority.

Growing up as they are in an ethnic enclave, then, most Tūrama School children’s experiences are of being a majority ethnic group. I quickly became aware of this when I was repeatedly asked ‘what culture are you?’ by children who had difficulty placing me within their emic ethnic classification system. I could not give any answer that satisfied them. While Māori and Pākehā are terms constituted in relation to each other—the hapū [subtribes] and iwi [tribes] of Aotearoa and the European explorers and settlers who arrived following James Cook’s voyage (Bell 2004)—these children did not hold a concept of my identity as Pākehā,

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3 Based on Education Review Office report. Many children at Tūrama School had both Māori and Pacific ancestry (or Māori and Pākehā) but as were designated as Māori.
let alone a sense of the colonial implications of my Pākehā status in relation to themselves. If I told them I was Pākehā, they were embarrassed; many of them used ‘Pākehā’ to refer to light-skinned Māori, but it carried a derogatory valence. If I said I was a New Zealander, they were confused. Some children thought that I was Māori, and one girl asked me if I was Samoan. Others decided that I was Scottish, English, or Irish. I came to realise that ‘culture,’ as these children are using it, indexes ancestry via country of origin, but constructed in such a way as to avoid the racial politics salient in their peer groups. I observed the concern with country of origin while observing the school clinic one day, when a boy who was waiting for his turn to see the nurse asks me, ‘what culture are you?’

‘Pākehā,’ I say.

‘No you’re not.’

‘Yeah, I am,’ I say, not all that surprised by this stage.

‘No, you’re Irish.’

‘I’m not Irish.’

‘You’re lying.’

‘I’m not lying.’

‘You are lying. You’re Irish. You have white skin. People with white skin are from… where are white people from?’

Based on the vague notion that lightly pigmented people originate from Ireland (or Scotland), children therefore explained my ‘culture’ as Irish or Scottish. While Pākehā New Zealanders typically conflate cultural and ethnic identity with nationality and emphasise country of birth—identifying as ‘New Zealander’ rather than British—Tūrama School children’s cultural classification system emphasises different core properties and applies the same logic to white people that white people would apply to everyone else. The children of Pacific or Asian ancestry identify, and are identified as Tongan, Samoan or Chinese, even when they or their parents were born in New Zealand. By that logic, white people should be identified by their ancestral origins, rather than their place of birth. Hence I, having mostly English ancestry, should identify as English.

I had an opportunity to expand my emerging sense of how Tūrama School children saw the world when I spent the day in Mrs Stevens’ class. Charismatic, opinionated, and with a uniquely laid-back teaching style, Mrs Stevens has a disdain for ‘political correctness’; she
does not hesitate to call children ‘retards’ or ‘juvenile delinquents’ and she describes her class as ‘special’ (insinuating ‘special needs’). Yet boys who rebel against their own teacher and are frequently disciplined for behavioural problems will come up to talk with Mrs Stevens. This year, all of the children who had truancy issues were placed into Mrs Stevens’ class so that they could be managed together, and so far, all of those children have been coming to school. Mrs Stevens jokes that she wouldn’t mind if some of them were truant more often, encouraging one exasperating boy to take a day off to give her a break.

For me, Mrs Stevens’ class often provided a window into the worlds of these children through her practice of promoting free and open conversation. She begins each day by discussing items in the news and this can prompt very long digressions; discussion of a news item about twins who were separated at birth ended up with a long laugh over photos of dogs that look like their owners. As part of these open discussions, children will make comments that reflect their worlds without the censorship that is usually maintained in the presence of adults. Children will share their knowledge of drugs: ‘I saw Mr. McPherson’s son smoking a lazy tinny’ [marijuana joint]; childbirth: ‘when you shit the baby out’ and gang affiliations: ‘Mozart is in a wolf gang aye Miss? Yeah like the Killer Beez.’

Mrs Stevens was also very happy to hold class discussions about issues of relevance to my work. When I expressed an interest in finding out more about the children’s perceptions of the ethnic breakdown of the country, we spontaneously decided to co-investigate this, integrating the teaching of percentages as the maths learning for the day. With the children lounging on the mat in front of the whiteboard, Mrs Stevens took them through a basic tutorial about percentages. Then I drew a long vertical rectangle on the board, divided into tenths and labeled from 0-100%.

‘So if this box is all of the kids at our school, where would I colour up to to show all of the children under 15 years old?’

‘You’d colour to the 100,’ the children correctly replied.

‘And if I wanted to show how many children are under 4 years old?’

‘Zero,’ they replied.

‘What about if I wanted to show how many are girls?’

‘About to the 50,’ they said.

‘Let’s do something a bit harder,’ I said, and Mrs Stevens stepped in.
‘How about how many children are Māori?’ she asked. This question elicited a range of answers, including 11-year-old Navahn’s facetious assertion that most children were Pākeha.

‘What are you talking about Navahn?’ Mrs Stevens waved her arms. ‘Do you see mostly Pākeha in here? There’s Chloe. That’s it.’

‘Who here is Māori?’ she continues. ‘Put your hand down, Joy, you’re Islander. Jasper, Trystan, put your hands up, you’re Māori. Paula…’ Mrs Stevens squints at the small girl in front of her. ‘I’m pretty sure you’re Māori.’

‘I’m Samoan,’ says Paula.

‘You can be both,’ I say. I am not sure why either of us are defining the children’s ethnicity for them.

‘So apart from an identity crisis, what we’ve worked out here is that about 70 percent of you are Māori,’ Mrs Stevens announces. She turns to me. ‘Which is about the schools average?’

‘Yeah, or just over, I think.’ I draw the line on the board in between the 70 and 75 and shade in the box, then add the shading for Pacific Islander (20%) and ‘other’ (Asian and Pākeha: 5%).

‘Ok so here’s a question’, I say. ‘What if we make this box every person in New Zealand. How many are girls?’ The class decides just over fifty percent—they have heard that women live longer than men. ‘Ok, so how much do you think we would colour in to show the number of people who are Māori?’

‘Ninety percent!’ A child shouts. ‘Seventy percent! Eighty! About fifty?’

‘Remembering that this school is 72% Māori,’ I say to the children who were calling out high numbers. ‘Ninety-five percent,’ they shout. Navahn guesses forty-five percent. I put marks along the box showing all the guesses. Most of them are above 75 percent. Navahn’s guess of 45 percent is by far the lowest.

‘Are you ready to find out who was closest?’ I ask. ‘I’m going to show you the answer.’ I hold my pen near the top of the box, and with dramatic effect gradually drop the pen lower and lower along the scale while the children gasp, to stop at around 15%. ‘This is how many people in New Zealand are Māori,’ I say. The children gape at me. I colour in the box to show Pākeha and other ethnic groups.

Seeing their astonishment, Mrs Stevens explains. ‘See the thing is, you guys live very insulated lives. Most of you have never even been out of Papakura! You look around you and think this is what everywhere looks like.’ She tells us the story of when she arrived in
Papakura and wondered, where are all the white people? ‘You think me and Julie and Chloe are the minorities, but actually we’re the majority in New Zealand!’

‘You guys are actually really rare,’ I say. ‘You’re special.’

‘Well, we already knew this class was special,’ Mrs Stevens jokes.

‘It’s funny,’ I comment to Mrs Stevens, ‘I suspected this was what they thought, but I also thought maybe—because they watch a lot of TV—that they would’ve picked up that Papakura looked different.’

‘But TV isn’t reality,’ she points out. ‘This…’ she gestures around the class. ‘Is reality.’

‘Shall we do one more question?’ I ask her. ‘Can we do a bit of a controversial one?’

‘You know me!’ Mrs Stevens replies. ‘Of course we can do a controversial one.’ I have done this exercise a couple of times in interviews with children, so I have an inkling of what to expect. I draw a horizontal line on the board, and mark it from one to ten. ‘On this line, ten is the richest people in New Zealand, and one is the poorest.’

‘Oooh,’ says Mrs Stevens.

‘Thinking about the kids who go to our school, on average, where do you think this school is?’

‘Nine!’ One child shouts. A range of numbers come in, most between four and nine. One girl chooses two, and Mrs Stevens nods at her. ‘You’re quite onto it today, aren’t you Joy?’

I have started to feel uncomfortable. ‘I haven’t quite thought this through,’ I mumble to Mrs Stevens. ‘Now we have to give them the answer.’

‘That’s okay,’ she brushes off my concern. ‘What is the answer?’

‘I was thinking school deciles,’ I said. ‘As a good proxy for the community.’

‘Oh yup,’ she says, and proceeds to explain the school decile rating system. ‘So every school is given a number based on things like how many people in the community are on benefits, how many people don’t have a job, how many people need state housing.’ I notice she is naming concrete things that would be familiar to these children. ‘If you don’t have many of these things in your community, then you’re a ten, and if you have lots, you’re a one. So, what do you guys think we are?’ The children’s responses do not really change from what was given previously.
Mrs Stevens circles the one, and the class falls quiet. ‘You guys are ones,’ she says. The children are wide eyed, blinking.

‘Well,’ I hasten to add, ‘schools that are ones still have some families who are rich, and schools that are tens have some families that are poor. And you guys actually have a really good school. What schools do you think get the most money from the government?’

‘The tens?’ Someone guesses.

‘No, the ones. I went to a ten school, and we didn’t have tablets.’

‘Yeah,’ Mrs Stevens adds, ‘we had to buy our own tablets for my kids. And you know what, you guys are ones now, but aim for a ten! Your parents are ones but you don’t have to be. Work hard, get your education, and go be tens!’

Despite this speech, the children are subdued, staring, processing. ‘They’re so quiet,’ I comment to Mrs Stevens and she laughs. ‘Yeah look, you’re all depressed now that we’ve burst your little bubbles.’

I wipe the graphs off the board and a child asks, ‘can you ask us more questions?’

‘Yeah,’ Mrs Stevens says with a wink at me. ‘Next time Julie’s in our class we’ll ask more questions.’

While being distinctly different from what might be considered ‘mainstream’ New Zealand, and representing the extreme end of the socio-economic scale in census data, what I took from this day was how normal this community and this school is from the perspective of the children living in it. This is despite the visibility of media discourses about child poverty and media representations of New Zealand lifestyles that are very different from the children’s own. This is the world for Tūrama School children, and this is the normal world.

Unequal childhoods

The struggles of Tūrama School families have until recently been largely invisibilised by the dominant cultural zeitgeist which reflects a nostalgic imagining of Aotearoa childhoods as happy, protected, resource-rich and world-leading (Tap 2007). According to the Child Poverty Monitor, 27% of New Zealand children are now considered to live in poverty, meaning they live in households with incomes less than 60% of the contemporary median after housing costs (Duncanson et al. 2017). International pressure has provoked serious
attention to the issue of childhood deprivation. As a signatory to the United Nations 2030 Agenda for sustainable development which came into effect in January 2016, New Zealand is required to halve the rates of 0-17 year-olds living in poverty from a 2015 baseline (United Nations 2015). Childhood inequity in Aotearoa has also been an increasing concern of researchers who document the short and long-term effects of poverty on health, education, and crime (Baker et al. 2012; Boston 2014; Fergusson, Horwood, and Boden 2008; Gibb, Fergusson, and Horwood 2012; Pearce and Dorling 2006; Rashbrooke 2013). This body of research includes three ongoing major longitudinal studies of child health: the Dunedin Multidisciplinary Health and Development Study, now on the second generation (Poulton, Moffitt, and Silva 2015; Silva 1990), the Christchurch Health and Development study of children born in 1977 (Fergusson, Boden, and Horwood 2015), and the Growing Up in New Zealand study which began in 2010 (Morton et al. 2013), all of which demonstrate links between early childhood environments and life trajectory. Another major longitudinal study of Pacific families in Aotearoa, initiated in 2000, investigates links between a host of developmental pathways and health incomes for Pasifika children, including culturally nuanced perspectives on the impacts of gambling, nutrition, alcohol consumption and violence on child health (Savila et al. 2011).

As child and adult morbidity and mortality statistics testify against the beloved cultural myth of an egalitarian society, action groups and the media have also drawn increasing attention to the numbers of children living below the poverty line, investigating, for example, inequalities in child educational achievement (Johnston 2015), housing conditions (Johnston and Knox 2017), and what children have for lunch (Barraclough 2017). Child poverty became a further issue in the 2017 election, and under pressure from the left-leaning opposition, during a debate the incumbent National leader made a surprise pledge to lift 100,000 children out of poverty within the next six years (Radio New Zealand 2017). This represented a major concession by a party who, up until that point, had disagreed with opposition parties, the Children’s Commissioner, researchers, experts and diverse organisations and action groups about the extent of and ability to measure child poverty in New Zealand (Peters and Besley 2014; Rashbrooke 2017). The struggles of impoverished adults, however, were rarely mentioned. In a neoliberal society where poverty is routinely discounted with discourses of individual responsibility (Boston and Chapple 2014), as the innocent and vulnerable faces of the future, children represent the one group who cannot be held responsible for their circumstances and thus headline appeals for policy change.
The children of Tūrama School are not immune from these discourses; they see the news, and so they, too, are concerned for these children living in poverty, perhaps in Africa, or with those homeless people they see on the street. Ten-year-old Amberlee does not like the Prime Minister John Key, she tells me, because he puts up the prices of the houses so the poor people can’t buy them. You can tell if someone is poor, she explains, because they don’t shave their face, and cart their things around in a trolley on the street, and poor children don’t have beds so they have to sleep on the floor in leaky houses. On a one-to-ten scale of poorest to richest, Amberlee guesses the children at Tūrama School are about a four, which means that on Monday and Tuesday you might bring lunch, but on Wednesday and Friday you have none. And some children, like this one girl, Helen, have no proper shoes and holes in their clothes.

Amberlee is particularly observant and her guess is one of the more accurate that children shared with me; whether I asked individuals, small groups, or entire classes like Mrs Stevens’, few rated Tūrama School less than five, and many thought they were a nine or a ten. Tūrama School straddles two suburbs which, according to the New Zealand Index of Multiple Deprivation, are in the 97th percentile for deprivation nationally, characterised by families with low median income, low frequency of educational qualifications, and high levels of government assistance benefits. Often, two or three families occupy the same household, like Ruby’s. Such statistics do obscure heterogeneity within the suburbs themselves, and some children at Tūrama School are considerably better off than others. However, Tūrama School is overall designated as a ‘decile one, step A’ school, indicating the highest of the 18 levels of socio-economic disadvantage used by the Ministry of Education to allocate funding.

Deprivation, in this context, manifests in some aspects of life more than others. Because state funding for Tūrama School is proportionately higher than for schools in more affluent areas, in some ways the children are quite well resourced. A cluster of around ten buildings, the school campus includes classroom blocks, a hall, several high-quality, safe playgrounds, sports fields and a swimming pool. Tūrama School children’s playtimes are enriched with play equipment and they have access to information technologies in the classroom and library, high quality teachers who care for and about them, smaller class sizes and several field trips a year for education or celebration. For some children, school represents the warmest, healthiest, and safest environment they inhabit, although for many, the classroom comes with its own sets of social challenges. Many children come to school inadequately
clothed and fed, and classrooms could be cold, leaky or prone to flooding. Those who came to their end of year graduation prize-giving wore a nice outfit, but half the graduating class were missing. Because parents often work multiple jobs with unconventional hours, their lack of availability means there are no school camps and no Saturday sports teams—both time-honoured traditions of New Zealand school life. Tūrama School children may still be developing their awareness of their inequitable circumstances, but their school experiences are still quite distinct from those at the decile ten school in the next district.

Tūrama School children are also not aware that in Aotearoa poverty has been found to increase the likelihood of a range of issues that impact children in the present and establish lifetime trajectories of disadvantage, including poor nutrition (Wham et al. 2015), health issues (Poulton et al. 2002), and educational achievement (Fergusson and Woodward 2000; Gibb, Fergusson, and Horwood 2012). Recent New Zealand research has demonstrated how the effects of poverty on brain development in the first few years of life can be carried throughout the life course. Results of the Dunedin longitudinal study show that around 20% of the population account for around 80% of adult economic burden, including 81 per cent of criminal convictions, 66 percent of welfare benefits, 78 percent of prescription fills and 40 percent of excess obese kilograms, and that these individuals, who were more likely to have grown up in socio-economic deprivation or have experienced child maltreatment, could be reliably predicted from neurological evaluation at age three (Caspi et al. 2016). Poverty is also linked to child abuse, which remains a significant problem in New Zealand. The White Paper on Vulnerable Children reports that between 7–10 children are killed each year by a caregiver, while in the year leading up to June 2012, the organisation responsible for care and protection issues, Child, Youth and Family (CYF, pronounced ‘sif/s’)⁴, found 4,766 cases of neglect, 3,249 cases of physical abuse, 1,396 cases of sexual abuse, and 12,114 cases of emotional abuse (often children who have witnessed family violence) (Ministry of Social Development 2012). Internationally, childhood abuse, along with other adverse childhood experiences including the loss, incarceration, mental illness, or drug/alcohol abuse of a parent, has been linked to a range of health conditions later in life including obesity, cancer, addiction, diabetes and stroke (Bynum et al. 2010; Felitti et al. 1998; Gilbert et al. 2015).

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⁴ Following numerous reports of present and historical mismanagement leading to abuse of children under CYF care, the agency was replaced by the controversial Ministry for Vulnerable Children (also known as Tamariki Ora) in 2017. Children, teachers and parents still commonly refer to the agency as CYFs.
These socio-economic pressures therefore have significant impact over the life course, and they create particular experiences of childhood. For Tūrama School children, their relative deprivation is mostly unseen, except perhaps for when they attend an inter-school kapa haka\(^5\) competition and notice the clean and fancy uniforms of the largely Pākehā private school also in attendance. This could be quite a different experience for socio-economically deprived children attending a high decile school. Yet Tūrama School children do notice who is requesting a stigmatised ‘spare lunch’, they manage their cold and sick bodies, and they are devastated by the poverty-related illness or death of family members. For many, family violence, or ‘getting a hiding’ is a normal part of life, but the threat of being ‘taken away’ by CYF means children will also work to conceal specific violent events from teachers.

These are the *practices* of children negotiating the structuring conditions of their lives: making meanings from material deprivation that in turn become the webs that govern their sociality; making accommodations in some domains to compensate for deprivation in others; caring for their health in the face of epidemic disease. The reasons for these circumstances, however, go much deeper, arising as a consequence of their colonisation and migration histories. In the following two sections, I outline the respective histories of these relationships with specific reference to Papakura and the health outcomes that disproportionately affect Māori and Pasifika children in this region.

**Colonial legacies and health**

The demographic pattern of Tūrama School, which shapes their views of the country, identities, and peer relations, has long historical roots which privilege Pākehā over Māori and other ethnic groups and which contribute to current health disparities in Papakura as well as nationally. The early colonial history of the Papakura region is well documented in several volumes (Craig 1982; Smith 2016) and by the Papakura Museum from archival material. I spent several hours in this museum building a picture of the events that lead to the current circumstances of Tūrama School children.

Post-colonial New Zealand history is built around an often fraught relationship between Māori and Pākehā which, since the signing of the Treaty of Waitangi in 1840, has seen Māori stripped of their lands, language and culture, marginalised, and set in motion cycles of inter-

\(^5\) Māori cultural performance
generational trauma which play out for the children of Tūrama School to this day. Craig (1982) details how the events following the Treaty established the Pākehā stronghold in the Papakura region. Prior to the Treaty, the South Auckland (Manukau) region appears to have been sparsely populated by local Māori, who roamed the wider area and resided seasonally in the Papakura area, taking advantage of a rich food bowl and cultivating some gardens. In the early 1840s three European families established a settlement in Papakura, and built what accounts suggest seem to have been friendly relationships with local Māori: sharing agricultural practices, settler children growing up speaking Te Reo [the Māori language] alongside their Māori playmates. This relationship became strained in the early 1860s. Under the Treaty, the crown had reserved a pre-emptive right to purchase Māori lands, and in 1853 the Governor, facing increased demand for land by settlers, reduced the price per acre, which saw an erosion of Māori holdings as the land was snapped up by wealthy speculators.

The colonial government then prepared to invade the Waikato region in 1863 to claim Māori land under the pretence of forcing the allegiance of local iwi to the Queen (Sinclair 2000). As the gateway to the Waikato region, Papakura was designated as a place of strategic importance, and a large military road was built connecting the two regions, which local Māori viewed with unease and protested. The tensions resulted in many Māori leaving the area to seek refuge in Waikato, while others were labelled as rebels, forced to leave the district, and large blocks of Māori-owned land, including villages and sacred places, were confiscated by the government, justified as punishment for a rebellion which had not eventuated. This land was parcelled up and distributed for settlement by military men who had fought in the Land Wars, and new European migrants arriving under the 1864 Waikato Immigration Scheme to settle the frontier between Auckland and the Waikato. From 1865 the Māori population of Papakura was all but eliminated, while descendants of the European settler families who developed the region into a thriving business centre remain in Papakura to this day.

The return of Māori to Papakura is less clearly documented, but from the late 1930s, population growth and the advent of World War Two opened up new opportunities for urban employment and resulted in a mass migration of Māori to urban centres. Between 1936 and 1986, an 83% rural Māori population became an 83% urban population (Belich 2001), many of whom tended to have less education and who, away from traditional whānau and hapū connections, struggled with poverty. Māori-run organisations such as the Māori Women’s Welfare League were set up to meet the needs of urban-based Māori, including housing and
pastoral care (Hill 2009 c.f. Gagne 2013). While the Māori cultural renaissance of the 1970s and 1980s saw the establishment of new marae in Auckland and the revival of Te Reo and Māori activism, many urban Māori did not hold a strong sense of Māori identity (van Meijl 2006; Borrell 2005). State housing was built to accommodate many of these Māori throughout Auckland. Meanwhile, Papakura saw the establishment of a military base in 1939, which brought considerable wealth and prestige to the area. The military base was closed in 1992, and the land was carved up for new developments, including state housing. In 2000, Papakura was again one of several areas of ‘high deprivation’ in South Auckland targeted for new state housing which became home to low-income Māori as well as Pacific migrants. The families of Māori children at Tūrama School, therefore, are from various iwi, and many maintain ties to marae in Northland, Waikato, or the rural East. Others have connections to the Papakura marae, which was established in 1979, and includes a G.P. and pharmacy service. Others still have little or no connection to Māori culture, may not know their iwi, and do not identify with a marae.

The price of colonisation has been high for Māori in general, who are disproportionately represented in statistics for almost every major indicator of health and wellbeing, from mortality and hospitalisation rates (well summarised by Reid, Taylor-Moore, and Varona 2014) to educational achievement (Marie, Fergusson, and Boden 2008). It should be noted that such statistics conceal the wide variation in circumstances across Māori from the top to the bottom of the socio-economic ladder (Durie 2001), and so the group of children I worked with at Tūrama School are not representative of Māori, but among the most disadvantaged of Māori. However these disparities can be consistently traced upstream not only to socio-economic disadvantage but also the intergenerational transmission of the historical trauma of colonisation, the collective disenfranchised grief, stress, and dislocation from culture that have destabilised the wairua [spirit] of whānau and iwi (Marie, Fergusson, and Boden 2008). These traumas are perpetuated through health care and education which is still, for the most part, delivered through colonising institutions, and Māori experience higher levels of racial discrimination which has documented negative effects on health and achievement (Blank, Houkamau, and Kingi 2016; Harris et al. 2006; Harris, Cormack, and Stanley 2013; Turner, Rubie-Davies, and Webber 2015). For the children I worked with, some steps have been taken to correct this, with a bilingual unit that operates semi-autonomously in Tūrama School according to Māori tikanga [customs], and a

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6 A central gathering place for Māori.
marae-based health service that several parents I spoke to used. New Zealand still has far to go before the damage may begin to be reversed, however, and the consequences of this colonial legacy can be seen in the health status of Tūrama School children, in particular, the rheumatic fever that caused Victor’s leaky heart valve.

Pasifika migration histories

Pasifika families in New Zealand, too, disproportionately share these strained socio-economic circumstances and subsequent impacts on health. Most of the 20% of children at Tūrama School who are Pasifika identify as Tongan or Samoan and were born in Auckland, though some are recent migrants. While historical processes of colonisation underlie the material disadvantage of Māori children, the inequitable living circumstances of Pasifika children stem from Tongan and Samoan migration histories. Large-scale migration from Polynesia to Aotearoa began after WW2 where migrants were encouraged as workers to help with labour-shortages in a booming economy. People from Tokelau, Niue and Cook Islands were deemed to be New Zealand citizens while migrants from Western Samoa and Tonga were required to apply for a work permit, though the practice of ‘turning a blind eye’ to those working beyond their work permit was commonplace as their labour was in demand (Savelio 2005).

From the mid-1970s, an economic downturn particularly affected manufacturing industries where many Pasifika were engaged and left communities struggling with high levels of unemployment. Societally, Pasifika peoples were cast as a threat to NZ and blamed for law and order problems (Dunsford et al. 2011). From 1974, ‘dawn raids’ were made on the homes of alleged overstayers and even some individuals with valid work permits were forcibly repatriated (Macpherson 1996; Liava’a 1998). These raids, often carried out at dawn to catch women and children while male family members did the night shift, created a climate of fear and made ‘islanders’ the targets of antagonism, stigmatisation and discrimination.

By 1991, the number of ‘Pacific’ people born in New Zealand equalled the number of migrants, and success in sports, arts and community work, including high-profile festivals, improved perceptions of Pasifika peoples (Dunsford et al. 2011). However, due to barriers such as language, education, and discrimination, Pasifika migrants and descendants are still often relegated into low-paying labour or shift-work and low-cost, poor-quality rental housing which adds stress onto families. Throughout the migration period until the present,
Pasifika peoples have struggled with housing, from discrimination by landlords and the prohibitive cost of renting or buying homes in Auckland. The radical policies of economic deregulation and liberalisation of the early 1990s led to a restructure of social welfare and corporatisation of state-housing programmes, including adopting of ‘market rates’ by Housing New Zealand which saw rents for low-income tenants rise by 106% between 1992-99 compared to a 23% increase in the private rental sector (Cheer, Kearns, and Murphy 2002). Consequently, housing related poverty became more widespread through the 1990s (Thorns 2000) and families often ‘doubled-up’ to save costs (Howden-Chapman et al. 2000), filling garages, sheds and caravans and in damp, uninsulated accommodation. As the government sold state-owned housing in gentrified, middle class neighbourhoods, rising rents saw the increased relocation of Pasifika—along with Māori—to outer, South and Western suburbs of Auckland (Dunsford et al. 2011), including Papakura.

Pasifika have also seen a shift in patterns of disease over the migration history, with the rise of non-communicable diseases such as diabetes and heart disease, along with lingering communicable diseases such as tuberculosis. For children, disparities in health measures between Pasifika and non-Pasifika in Aotearoa have been apparent since systematic tracking of Pasifika children began in the 1970s (Dunsford et al. 2011), including obesity (Anderson, Gorman, and Lines 1977; Bell and Parnell 1996), nutritional deficiencies (Bell and Parnell 1996; Wilson, Grant, and Wall 1999), asthma (Ellison-Loschmann et al. 2009; Mitchell and Cutler 1984) and communicable diseases including measles (Hardy, Lennon, and Mitchell 1987; Steele 1973) meningococcal A and B (Baker et al. 2000; Lennon et al. 1993; Wilson et al. 1995), and respiratory illness (Grant 1999). Of particular interest to this thesis, disparities for rheumatic fever have been recorded since the 1980s, and from 1980–1984 the rate for Pasifika children under 15 years old was between 9–10 times the rate from European children (Lennon et al. 1988). This discrepancy has continued to this day; in 2016, after four years of intervention programmes rheumatic fever rates for Pasifika children are still nearly four times higher than for Māori and eight times higher than the national rate (Ministry of Health 2017). For many schools in low socio-economic areas of the country, efforts to combat rheumatic fever have become a prominent feature of school life, meaning that even though only a handful of children at the school would ever develop the disease, it has become significant to the coproduction of health for every child at Tūrama School.
Rheumatic fever

Among the confluence of physical and social disruptions that have marked Victor’s body, it is the heart valve damage following rheumatic fever that has attracted recent government attention and intervention on a national scale. An autoimmune response to group A streptococcus (GAS) infection, acute rheumatic fever is a major cause of childhood morbidity and mortality in mainly developing countries (Carapetis et al. 2005). Rheumatic fever occurs mainly in children aged 5–14 with mean incidence in New Zealand peaking at age 9–12 (Milne et al. 2012).

The chief concern is the inflammation of the heart (carditis), which can cause damage to the cardiac valve or ‘heart damage’ as it is described in lay terms (Jaine 2008), and can lead to chronic rheumatic heart disease (CRHD) in later life (Milne 2012). While rheumatic fever is rarely fatal for New Zealand children, the CRHD that often follows is a significant cause of premature death in Aotearoa, and the State’s failure to prevent rheumatic fever now means a burden for decades to come for those communities worst affected (Wilson 2010). The experience of a common childhood GAS infection—strep throat—can therefore result in a sequelae of subsequent illnesses for the rest of an individual’s life.

<table>
<thead>
<tr>
<th></th>
<th>Overall mean incidence rate</th>
<th>Increase from 1993-2009</th>
<th>Mean incidence rate for decile 9/10 (highest deprivation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Māori</td>
<td>40.2</td>
<td>79% increase</td>
<td>64.9</td>
</tr>
<tr>
<td>Pacific</td>
<td>81.2</td>
<td>73% increase</td>
<td>96.0</td>
</tr>
<tr>
<td>Non-Māori/Pacific</td>
<td>2.1</td>
<td>71% decline</td>
<td>&lt;10</td>
</tr>
<tr>
<td>All children</td>
<td>17.2</td>
<td>59% increase</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Mean rheumatic fever incidence rates per 100,000 from 2000-2009 for children aged 5-14 in New Zealand. Data from Milne et al., 2012.

Rates of rheumatic fever, a disease associated with developing countries, increased in New Zealand by 59% between 1993 and 2009 (Milne et al. 2012) (Table 1), driven almost exclusively by rising incidences for Māori and Pasifika children (Jaine, Baker, and Venugopal 2008). The discrepancy is startling; mean incidence rates between 2000–2009 were 81.2 (per 100,000) for Pasifika children and 40.2 for Māori children, compared to just 2.1 for non-Māori/ Pasifika (predominantly Pākehā) children. A 2012 epidemiological study found that while Māori and Pacific children comprised 30% of children in the 2006 census, they accounted for 92% of new cases of rheumatic fever in the period between 2000–2009.
Although twin studies indicate a genetic susceptibility to rheumatic fever exists at an individual level, there is no evidence that these ethnic disparities reflect population-level genetic differences (Bryant et al. 2009; Carapetis, Currie, and Mathews 2000). Even in populations heavily exposed to GAS bacteria, no more than around 3–6% of children are expected to develop rheumatic fever (Carapetis, Currie, and Mathews 2000).

![Graph showing rheumatic fever incidence rate by socioeconomic deprivation](image)

**Figure 1.** Rheumatic fever incidence rate by socioeconomic deprivation (mean of 2000-2009) for Pacific (green), Māori (red), and non-Māori/Pacific (blue). Reproduced from: Milne et al. 2012.

The overrepresentation of Māori and Pasifika children in New Zealand’s rheumatic fever statistics is therefore the result of a complex interplay between biological and socially mediated factors: individual susceptibility and GAS virulence, and levels of bacteria exposure and treatment with antibiotics (Carapetis, Currie, and Mathews 2000). These factors appear to be influenced by socio-economic disadvantage (household overcrowding, increased incidence of streptococcal upper respiratory tract infections, decreased health care access) (Jaine, Baker, and Venugopal 2011; Milne et al. 2012; Wilson 2010). A recent epidemiological study in New Zealand demonstrated a steep increase in incidence rates for rheumatic fever with degree of socioeconomic deprivation (Milne et al. 2012), with the highest deprivation quintile accounting for 70% of hospital admissions among children aged 5-14 (Figure 1). The cumulative risk for a child living in a region of high socio-economic disadvantage—like the area where Tūrama School is located—is 0.0066; about a one-in-150
risk of being admitted to hospital for rheumatic fever (Milne et al. 2012). At Tūrama School, this child is Victor.

The experience of rheumatic fever for children like Victor begins with a lengthy hospital stay followed by ten years of monthly prophylactic penicillin injections to prevent secondary recurrences. Victor, like most children, receives his injection at school, where the district nurse comes by to administer the injection to him and several others. However, after incidence rates in these socio-economically disadvantaged groups rose above a threshold of >50 per 100,000 children, attention shifted to primary prevention. In 2011, the New Zealand government established the Rheumatic Fever Prevention Programme (RFPP) with the aim of reducing rheumatic fever admissions by two thirds (Ministry of Health 2013). The major component of the RFPP targets GAS bacteria, with funding allocated for free throat swabs for high-risk groups at primary care rapid response clinics and free clinics in schools in areas of high economic-deprivation. A secondary component involves health promotion to families and health professionals. Least emphasis is put on the third aspect: strategies to address social and environmental factors such as poor housing and household crowding. Funding was also allocated for research, surveillance and primary care development.

Thus, the disease which has shaped Victor’s childhood is now structurally integrated into the lives of all children at Tūrama School through their ‘sore throat clinic’, which links Victor to children like Dante, who visits the school clinic almost weekly to have his throat checked, and Teuila, who diligently takes her antibiotics only every second day so that her siblings might have the other half, and Marielle, who tells her friends she has rheumatic fever, not realising her diagnosis is actually for strep throat. While the clinical and political implications of rheumatic fever policy are to greater and lesser degrees the subject of policy attention, what lies underneath is the power of the clinic to infiltrate, just as it is infiltrated by the institutional activities that bring it to life in situ. The clinic becomes woven into classroom dynamics while simultaneously restructuring the school day; it is both synthesised into new productions of knowledge and reconfigures children’s relationships to their bodies. Chapters four and six describe the processes through which this occurs, as underpinned by the roles of the state and the institution in child health detailed in the next section.
Ruby had been experiencing stomach pains for several days but remained at school while the office and her teacher, Mrs Randall, tried unsuccessfully to contact her mother. ‘You need to tell your mum to take you to the doctor,’ Mrs Randall told her, stern with worry, as Ruby curled up against me, hot and listless. The next day Ruby did not come to school, but she was back the day after, swinging on her chair. ‘Are you feeling better?’ I asked, and she nodded. ‘Not really,’ said Mrs Randall, speaking for Ruby. Later that day, she tells me she’d asked Ruby whether she went to a doctor. ‘She said she had,’ said Mrs Randall, ‘and I asked her what the doctor said, and she said the doctor said she had a sore stomach. So I asked if the doctor had checked for appendicitis and she said yes. But I don’t think she went to the doctor at all.’ Mrs Randall thought Ruby was covering for her parents. The family had access to a doctor at the marae, so Mrs Randall reckoned Ruby’s parents ‘couldn’t be bothered’ taking her. ‘But I will keep calling them and telling them she needs to go,’ Mrs Randall concluded.

Much responsibility for managing the problem of children in poverty ultimately falls on New Zealand schools, but the multiple roles played by the institution in caring for children’s health and wellbeing sit in tension with neoliberal discourses that frame child poverty as a problem of parental responsibility. Situated as it is in an economically deprived area, Tūrama School participates in an especially large number of such initiatives, supported by national or local government or non-government organisations, for the purpose of caring for children’s bodies. At the time of my fieldwork, these included government-funded services (the dental service, fruit in schools, throat and skin checks, social workers in schools); corporate-sponsored services: (milk in schools; school provided services: basic first aid, some provision of school uniform, spare lunches, sports equipment and lunch time sports games); and NGO sponsored services: (head lice checks and treatment, breakfast in schools, visits from the Life Education bus, provision of jackets, shoes and basic hygiene supplies [tissues, hand sanitiser]). The school itself supports all of these initiatives with class time, organisation, distribution and administration. As well as these formal initiatives, the school has an informal role in looking after children’s bodies, and teachers and other staff regularly diagnose and make decisions on children’s ailments, and actively educate about or promote healthy behaviours. There are also external services, such as CYF, which often operate through the school.

For Tūrama School, this creates an environment where teachers grapple with the boundaries of their role, alternately motivated by a desire to ‘make a difference’ and frustrated and
fatigued by the demands of caring for twenty-odd children while under Ministry of Education pressure to improve student performance in the national standardised tests. Teachers generally made their own decisions about what role to play, and so Mrs Charles gave Ruby and other pupils her phone number and picked up Mila one night from the train station in the next district after she ran away from home. Mrs Randall intervened to make sure Ruby saw a doctor and used her own money to buy stationery for one girl and a swimsuit for another so that she could join the class in the pool. Mrs Stevens, who drives one boy to school every day to ensure he actually attends, reflected that while these things should be the parent’s responsibility, in her view, schools had to be ‘realistic’ and that, for instance, when parents are not taking children to a doctor, the school needs to step in. ‘Shouldn’t have to, but I think they do have to,’ she concluded.

This conclusion, we shouldn’t have to but we do have to, and the accompanying fatigue and resentment, is shared by many of the staff, and the cause of great internal conflict. A senior staff member, frustrated by the intrusion of health and NGO services into education, sums up, and then immediately criticises his own position: ‘Me, I would say no milk, no fruit, no free jackets, no shoes, which is probably not a very good decision.’

The role of state schools in child health is not new, but understandings of the role have shifted and to a large extent, become more ambiguous under neoliberalism. Aotearoa has an established history of embracing a ‘welfare state’ for population health and wellbeing, and with particular concern for children. The first decades of the 20th century saw an emerging interest of central government in child health and welfare, beginning with a raft of policies legislating for lower income-tax for families with dependants (1914), a minimum-wage calculated for the support of children (1920), and a benefit for families with more than two children (1927) (Easton 1980). The welfare state was entrenched in the 1938 Social Security Act which encompassed a range of material benefits for the protection of particular groups, including widows, the disabled and the elderly, followed by the 1946 universal family benefit, and the domestic purposes benefit for single parents in 1973. Throughout this time, schools have functioned as key sites of state health surveillance and intervention (Burrows and Wright 2007; Kearns and Collins 2000). From as early as 1920, school dental clinics were established to supply free dental treatment to all children until age 18 (Saunders 1964), while health and hygiene was a key concern of early New Zealand industrial schools for homeless or delinquent children (Matthews 2000). By the late 1940s all children underwent routinised medical examinations at school and those who were not ‘thriving’ in their home
environment were sent to state-sponsored ‘health’ camps (Kearns and Collins 2000), some of which persist today, though there are far fewer and they cater to a much narrower group of children. Aspects of state care continue in the present, including publicly funded health services, and a 2004 Working for Families benefit which supplements the incomes of working parents (Taxation (Working for Families) Act 2004).

However, the 1980s and 1990s saw dramatic shifts in economic policy and state welfare, the effects of which on low-income populations are still felt to this day (Poata-Smith 2013). The once-popular ethos of the welfare state eroded from the mid-1970s as economic and political changes led to rising inflation, public debt and unemployment. In 1984 the Labour government adopted the economic policies of the early 1980s U.K. and U.S., known as ‘neoliberalism’, to dramatically restructure state functioning in accordance with competitive market principles: opening up the economy to international markets and corporatizing and privatising state-owned commercial operations and assets (Larner 1997). The effect of these was a further rise in unemployment, peaking in 1991. This peak coincided with the introduction of major welfare reforms by the new National government, which slashed benefits, restructured state housing provisions to require profit, and introduced employment law reform to deregulate the labour market, remove protections for workers, and undermine collective wage bargaining (Larner 1997; Nairn, Higgins, and Sligo 2012). The subsequent rise in poverty disproportionately impacted Māori and Pasifika families, who were more likely to be employed in low wage manufacturing or on benefits [welfare], and particularly impacted children, of whom 29% were in the lowest income quintile (Atwool 1999; Poata-Smith 2013). By 1998, social policy initiatives were being discursively framed in terms of ‘social responsibility’, materialising in the distribution throughout the New Zealand population of a pamphlet entitled *Towards a Code of Social & Family Responsibility, a Public Discussion Document*, which stated an aim of clarifying the responsibilities of families, particularly parents, in order to reduce government spending (Larner 2000; Tap 2007). Further welfare reforms in 1998, 2006 and 2013 continued to tighten the criteria for receiving a benefit and reconfigured welfare payments in terms of employment-seeking support (Social Security Act 1964). Heavy penalties for not meeting benefit criteria were reported to have cut payments for 43,000 parents in the two years following the 2012 reform, alarming child welfare interest groups (Migone 2015).

The rise of neoliberalism also saw a withdrawal of central government from schools, and a reframing of schools as ‘self-managing’ under a 1989 reform (Wylie 1994). The idea was to
move away from a one-size fits all approach to child health and hand over the power and decision making process to local communities to decide what the priorities are for their community, and what approach is most useful. Although Sinkinson (2011) notes that educating individuals to be accountable for their own health status has been an enduring aspect of the curriculum since the early 20th century, at the same time, neoliberal ideologies infused schools with ‘healthism’, discourses of health based on an ideology of individualism and presupposing that individuals should take responsibility for their own health status. For example, as part of the ‘Health Promoting Schools’ movement initiated by the World Health Organisation, the 2005 Fruit in Schools programme gave a free piece of fruit to each child in low decile schools, but it came tagged with an educational mandate to ‘promote healthy lifestyles’ and approaches that ‘empowered’ students with the responsibility for educating their peers and creating health promoting programmes and environments (Boyd 2011).

However, in localising the solutions, the state also localises responsibility for the problems; problems that in many cases are produced by structural conditions much greater than the local community. Children’s dietary habits are constrained by the affordability of fruit, itself governed by national and global markets, seasonal climates, pests and pathogens, as well as national-level policy factors, including tightened restrictions on access to benefits, increasing rents from a housing shortage in Auckland city, a low minimum-wage level, and epidemics of addiction, domestic violence and mental illness (Reid, Taylor-Moore, and Varona 2014). While Tūrama School children largely conceptualised health in terms of eating fruit and vegetables, for many, the government provisions were the only fruit they could eat that week. The intent to establish responsibility for healthy eating without changing the constraining conditions means that while technically a Health Promoting School, Tūrama School has not followed the criteria for the last few years, and the only aspect of the Fruit in Schools programme remaining is the free fruit.

The economic and housing insecurity wrought by neoliberalism also contribute to high rates of transience as families move when rents rise or children are passed between kin. Polynesian childhoods are embedded in the wider family, whether that be the Māori whānau, the Samoan aiga, or the Tongan kainga, and within this fluid kinship, adult kin other than parents are typically seen as equally influential in the support and guidance of children (Brady 1976; Carroll 1970; Metge 1995; Morton 1996; Ritchie and Ritchie 1979). Widespread across Polynesia are ‘whāngai’ practices, which loosely means adoption (but not in the formal or legal sense of the English word), where children are placed with kin other than biological
parents for reasons such as continuing a family name, strengthening whānau bonds, or alleviating financial pressure. Though the introduction of the Domestic Purposes Benefit in 1973 meant that single mothers could now care for their children themselves (Metge 1995), in the present day, children can still commonly be ‘whāngai-ed’ out to other relatives, particularly grandparents, in the short or long term when circumstances—especially economic and housing instability—necessitate (Gagné 2013). Thus Ruby’s youngest brother is also her auntie’s son, and both of them live in Ruby’s household. Often, when children are in situations of abuse or neglect, whānau members, particularly kaumātua [elders], may intervene and negotiate for the child to stay with another whānau member. Hence nine-year-old Whetu lived with his grandmother for several years, and ten-year-old Cassidee and eleven-year-old Trystan lived with their respective aunties. In general, when CYF ‘uplifts’ (their terminology) a child for safety reasons, they will try to place the child with whānau, though this is not always possible. Many Tūrama School children therefore move back and forth between kin, often changing schools as well. New Zealand family policy and education systems, however, are based on Western beliefs and values, including an assumption that the nuclear family is the main, and superior unit of social organisation (Durie-Hall and Metge 1992; Metge 1995; Morehu 2005; Ritchie and Ritchie 1979, 1997). This means that the basic design of policies for housing, health, employment, education and social services, as well as legal concepts of adoption, guardianship and parental rights, have not always effectively taken into account the cultural needs of the people who could most benefit from the support. One way in which this issue impacts children is that the institution-based education systems are designed for stable, nuclear families and do not accommodate the transience of whānau or the many children who ‘boomerang’ (Gilbert 2005: viii) in and out of schools such as Tūrama School over the year. For children, as I discuss in chapter seven, shifting schools during the school year can mean disrupted peer and teacher relationships and impact academic progress (Gilbert 2005; Mutch, Rarere, and Stratford 2011; Wynd 2014), while the social instability that this transience creates has important implications for health. Children’s health, therefore, is profoundly shaped by the realisation of neoliberal ideologies through the school institution: in the formal interventions that offer throat checks and free fruit; in the way that teachers balance finding the boundaries of their role with their own affective sense of care; or in the multiple transitions pupils must make from one institution to another. But children do not passively experience the impacts of these structures on their health; they actively meet, negotiate, and transform them. What remains missing from this
picture is what I have alluded to throughout this chapter: the experiences, understandings, and collective meanings and practices of children themselves which contribute to the coproduction of these environments, their bodies, and their health. The remainder of this thesis will therefore forefront children’s perspectives of living and practicing these Aotearoa childhoods.
Chapter Three

Negotiating Generational Differences in Ethical Research

Two interacting but distinct worlds coexist within Tūrama School; the sparsely populated but hierarchically privileged adult world, and the populous but subordinate world of children. These worlds are maintained by separate sets of written and unwritten rules for staff and students in the school, but also the organisation and daily practices of members of both. Space, which appears to be shared, is fractured with tiny implicit boundaries demarcating who goes into which place and what they do there. In the classroom, children occupy small desks accompanied by small chairs, and sit or lie on the mat, while teachers have big desks and sit on big chairs at the edges of children’s space. Elsewhere, adults congregate in staff-only spaces, offices and the tea room, while children requesting help or delivering a message will stand with their toes in line with the door frames, cautiously peering in, waiting to catch the attention of an adult occupant. Children’s habitats are found in the familiarity of the playgrounds and sports fields and the school library, while teachers patrol the peripheries in fluorescent safety vests. In assemblies, children sit in rows on the hard hall floor, surveyed by teachers who sit at the end. Their spaces are further partitioned by age and gender, and these are maintained by children themselves; seniors guide wayward juniors back over the line painted through the middle of the hot asphalt courts while shrieks caution boys to steer clear of the girls’ bathrooms. The school is patterned by invisible stripes and spots, zoned by identity and status, marked by boundaries that are not seen but felt, the sense of exposure as eyes are drawn to a body that is disproportionate to those around it, out of place. This architectural and physical segregation of the child and adult populations is what Foucault (1979) identifies as a mechanism of power which works to enable access to the bodies, actions and attitudes of individuals. Adults hold absolute authority here, but not absolute power, as children adapt to, ignore, resist or internalise power structures in their own quiet or unquiet way.

The school’s architectural ordering, its temporal organisation by age group and time of day and season, and its governing rules and systems are adult creations to maintain adult authority (Foucault 1979). Within this space however, to use Corsaro’s analogy, children work like
spiders, spinning webs of meaning across everything they find (Corsaro 2015). These webs are largely disregarded by adults, if they are seen at all, and coming to understand children’s cultures, without disrupting, misrepresenting, or colonising them has been a problem of adult researchers for over three decades. In this chapter I consider one aspect of this problem: the issue of navigating generational differences in fieldwork. Although focussing on research ethics, this chapter also foregrounds the relational in producing the knowledge which supports this thesis, and thus sets out the epistemology of what is to come.

It was into this highly segregated world that I arrived, an adult most similar not only generationally but ethnically and socioeconomically to the authoritative adults, rather than the children from whom I wanted to learn1. Anthropologists and sociologists (Christensen 2004; Fine and Sandstrom 1988; James 2001; Mandell 1988; Mayall 2000) have long remarked on the role of the researcher as a central, and uniquely challenging issue in studies with children, and this is particularly the case in institutional settings such as schools where differences in status and authority are most systematically, ritually, and symbolically drawn and maintained. In anthropology the role of the researcher—or who the researcher is in relation to others in the field—has been vigorously debated with regards to adults (Adler and Adler 1987; Gold 1958), particularly with the rise of engaged or activist anthropology (Scheper-Hughes 1995), and is certainly also a source of ethical or practical tension in adult research (Kloos 1969). However, the relations of authority embedded in child and adult bodies create particular problems in fieldwork with children. In studies where children are accompanied by adults, it is almost inevitable that adult concerns are asserted. Moreover, subgroups such as parents and teachers both hold authority over the children who are the focus of research, and act as gatekeepers regulating researcher access and child participation. Finally, in traditional ethnographic settings researchers tend to assume an equal status with their research subjects—or at least that participants should be treated as such. Fine and Sandstrum (1988), however, argue that child research is distinct even from other protected groups such as the mentally disabled, because the social code requires that legitimate adult-child relationships be governed by adult authority.

1 Differences in ethnic and cultural background, class, and from half of them, gender, posed various limitations to accessing and building relationships with my participants. However, it was the generational differences—specifically the intersection of these other identity categories with generational differences—that created relational tensions in the field. While other teacher aides were from the local Māori community and parents of children at the school, I was ethnically marked as aligning with the Pākehā senior teachers and management.
Nevertheless, the focus of ethical guidelines for children tends to be on entry to the field and discrete ethical moments (Christensen and Prout 2002)—consent, confidentiality, protection from harm—rather than on the ongoing, performative practice of intersubjectivity that constitutes fieldwork (Meloni, Vanthuyne, and Rousseau 2015; Shore and Trnka 2013). For adult researchers working with children, bypassing the authority structures embedded in generational relations without undermining them, and navigating the expectations of other adults while maintaining allegiance to child participants is an often impossible task. This is a particular problem with participant observation, where the researcher hopes to integrate themselves into a natural setting and must work within established power structures, in contrast to interviews, focus groups or workshops which are already marked as an out of the ordinary event and may be constituted within a special set of rules.

With all these complications, how then are adult researchers to approach participant observation based fieldwork in schools? In this chapter I consider this question in reference to popular suggested researcher roles, namely the ‘friend’ (Fine and Sandstrom 1988), the ‘least adult’ (Mandell 1988) and the ‘unusual’ or ‘different sort of adult’ (Christensen 2004). Though psychological paradigms have tended towards more detached observational roles, the social sciences have been concerned with understanding data as produced through relationships, hence researchers such as Mandell (1988), Mayall (2000) and Christensen (2004) seek roles that explicitly avoid authority, but facilitate acceptance by children as participants in their social interactions. While these researchers acknowledge that there is no one ideal role of the adult researcher, nor is any role free from tensions and dilemmas, I discuss two specific problems I encountered within my ‘different kind of adult’ role. Firstly, there can often be a discrepancy between how adults expect or perceive children to be, and how children actually are. Researchers who are granted access to children’s worlds can become mired in political ‘stickiness’ when confronted with children’s taboo behaviours, caught between conflicting ethical and cultural values. Secondly, children rely on known categories of adult ‘type’ in order to successfully navigate the political quagmire that is being a child in an adult dominated institution. While I was concerned with building rapport, in trying to distance myself from adult authority, the ambiguity of my role from children’s perspectives created the conditions for disclosures which required me to break confidentiality, with significant impact for participants.

I therefore highlight the importance of relational perspectives on researcher roles, including the perspectives of adult and child participants who must read and interpret this role in the
context of their daily lives. To navigate the tensions that can arise from intergenerational fieldwork, I propose the ‘transparent guest’ as an operational mode for fieldwork. Although similar to the ‘different kind of adult’ role, the transparent guest has two main features which help to demystify how intergenerational structures and rules are negotiated. Firstly, most children have either hosted, or themselves been a ‘guest’, and understand that guests can bring with them rules from home which are different to those in the host environment. The role of ‘guest’ therefore establishes the researcher as someone governed by a set of rules different to either adults or children in the school. Secondly, a policy of transparency, of naming what the researcher can or cannot do and why, makes those rules visible to children and other adults in the institution and can help to minimise issues arising from unknown or unpredictable codes of behaviour. Researchers may then also deal with politically sticky situations by enlisting the help of children sympathetic to the worry of breaking a rule or getting into trouble. Importantly, both of these aspects form part of an ongoing, relational and co-constructed process of negotiating how researchers and participants can be with each other.

Roles of the adult fieldworker in research with children

Accompanying the burgeoning interest in child-centred research over the last three decades, the recognition that children have their own cultures (Corsaro 1979, 1992; Opie and Opie 1969), the turn towards research ‘with’ rather than ‘on’ children, and the children’s-rights framework have all implicated the taken-for-granted role of adult researcher as critical to research ethics and validity. A number of researcher roles have been proposed in variable attempts to minimise or mitigate the power differences between child participants and adult researchers, and allow adults greatest access to children’s private worlds while maintaining protections with what is seen as a uniquely vulnerable group. This is particularly a problem of the Western world, where children are cordoned off into separate, ‘child-friendly’ spaces and there are few opportunities for children to interact with adults outside of the authority structures of parent-offspring and teacher-student (James and Prout 1990).

One solution has been to engage children as researchers, bypassing the problem of generational differences completely, or as research partners who facilitate the research process with their peers. This approach can, however, mean shifting the problem one step upstream, as researchers still need to negotiate their relationships with child partners. Others
have argued that child-led research is not necessarily ‘better’ or more ethical (Gallacher and Gallagher 2008), and certainly there are practical obstacles to child-led research. My own attempts to involve children as active partners were rather unsuccessful; I invited children to share their thoughts on my research questions or findings, but most children would shrug and say they didn’t know. Realising a partnership would require giving children greater understanding of research purposes and methods, I began devising a set of workshops modelled after the child-led work coming from Children’s Research Centre at the Open University (The Open University n.d.), with the idea of producing a book based on children’s own ideas and experiences. In the end, this idea proved too ambitious for my limited resources and the institutional constraints of the school. Instead, I brought paper and pens to the outdoor table or to the library at lunchtimes and any child could come and draw with me. I did not tell them what to draw, but I said I was making a book about them, and invited them to contribute by telling me what should be in the book or drawing a picture of either something they really liked to do, or a time when things were difficult. Children also drew during interviews with me and in class time when they had finished their set tasks. In addition, I worked with small groups of children, recording their conversations about ‘what it is like to be a child’. This was a topic they took to with gusto. While they drew, I compiled their thoughts and illustrations, along with some of my own, into a picture book on my laptop, adding photographic backgrounds and digitally colouring the pictures [see appendix]. Children would come, peer over my shoulder, and ask questions but when I invited their thoughts they had little specific feedback, beyond nodding or saying it was cool. ‘We do do that,’ sometimes they would exclaim in surprise.

My attempts at explicit partnership, therefore, were rather unsuccessful, at least in part because, embedded as we were in everyday school contexts, mutuality could gain little traction when the institutionalised relational mode was hierarchical. Regardless, in many cases, adult researchers still wish to engage with child participants directly, and I do not think good research with children necessitates their partnership as coinvestigators. When working with children, then, what role should the researcher assume?

In her discussion of this question, Nancy Mandell (1988) classifies proposed researcher roles into discrete categories of participation level from detached observer to complete involvement. Alternatively, these roles can be seen as falling across a spectrum which reflects the degree to which the researcher accepts or rejects the social implications of their own adulthood (Table 2). At one end is what Mandell considers the essentialising position of
deterministic fields, for example developmental psychology, which hold that children’s worlds are so distinct as to preclude adult participation of any sort, and the structure of age roles and adult ethnocentrism immutable. At the other end, researchers such as Mandell (1988), Waksler (1986) and Goode (1986) contend that differences between adults and children are mostly ideological, and therefore all aspects of adultness except for physical size may be cast aside. Child researchers in social science tend to propose roles which fall between the middle and this latter end of the spectrum, including the role of friend (Fine and Sandstrom 1988), least adult (Mandell 1988), and unusual adult (Christensen 2004; Mayall 2000), all of which accept to some degree the identity of adult, but attempt to disrobe from the conventional authority and ethnocentrism usually associated with adults.

<table>
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<th>Adultness immutable</th>
<th>Adult researchers cannot access children’s worlds</th>
<th>Adult researchers can access children’s worlds as a different of unusual kind of adult</th>
<th>Adult researchers can access children’s worlds as friends</th>
<th>Adult researchers can participate as children or least adult in children’s worlds</th>
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Table 2. Spectrum of adult researcher roles ranging from a view of adultness as immutable and precluding any access to children’s cultures, and almost complete rejection of adultness apart from physical differences.

Towards the right of the spectrum, Fine and Sandstrom (1988) propose viewing roles as functions of two axes, with the roles they identify—supervisor, leader, observer, and friend—representing four permutations of degree of authority and positive relation. The authors advocate the latter ‘friend’ role as a semi-participatory approach, seeing its low authority and high positive relation as most conducive to ethical research. However, adult authority is often not easily dismantled, and in many settings, such as institutions, researchers may be limited by the requirement of other adults that they align with more adult roles. The ‘friend’ role also does not address the question of how generational differences may be approached, especially given the social illegitimacy of adult-child
friendships in a culture where the proper mode of adult-child relations involves the authoritative and responsible adult who protects and educates children.

Further still to the right of the spectrum, Mandell (1988) posits the ‘least adult’ role, which, though criticised, still holds a great deal of influence (e.g. Randall 2012). The role involves full participation in children’s activity in the ‘least adult manner possible’. In her embodiment of this role in her fieldwork with 2-4 year-olds in two day-care centres, Mandell initially observed, then imitated children’s actions, oral expressions, and social exchanges, through trial and error coming to be, she claims, accepted as a kind of big child. By ‘following them into the sandbox’ (1988:45) Mandell became included in children’s play and through experience learnt the rituals and rules children create and use to govern their interactions. (Similar methodology was used to the same effect by Corsaro (1979), but Mandell takes this a step further from peripheral to full participation). In attempting to minimise all difference between herself and the children, Mandell did not direct or correct children’s actions, even participating in rule breaking or stretching on occasions, and avoided adult ‘helper’ roles where possible.

Mandell’s ‘least child’ role has been critiqued by Mayall (2000) and Christensen (2004), both of whom arrived at an approach of ‘different’ or ‘unusual kind of adult’ which sees accepting the adultness of the researcher as important for understanding children’s experiences, but attempts to create a new, different category of adult. While the ‘least child’ approach attempts to show children that differences between adults and children can be diffused to near invisibility, Mayall argues that to children, a central characteristic of adults is that they have power over children. Good data on children’s experience therefore cannot begin with downplaying that truth. Instead, Mayall’s approach is to work with generational issues: ‘asking children directly, to help me, an adult, to understand childhood’ (2000:122). Through this approach, Mayall became recognised as a ‘non-official adult’ in school settings, a person to whom children could complain about school, but still relate to as an adult. Moreover, in facilitating focus group conversations (an adult role) but unlike adults, sitting back from directing the discussion, Mayall was able to listen to the way that children used conversation to acquire—and produce—knowledge. By shifting her own behaviour, Mayall found that children started to set aside the generational order and move out of ‘adult-question, child-reply mode’ (2000:126). Meanwhile, it was sometimes only through augmenting children’s voices with parent and teacher accounts that Mayall could fully understand a child’s experience.
Christensen (2004), meanwhile, criticises the least adult role as idealistic ‘wishing away’ differences between children and adults. Like Mayall, her approach in fieldwork in a Danish school context was to embrace her status as adult, but consider with greater nuance the question ‘what is an adult’, recognising that when children ask the ubiquitous ‘who are you?’, they are engaging with wider notions of who we are to each other. What kind of adult the researcher embodies very much matters in determining the kind of relationships established and the kind of data that is elicited, and how ethically these interactions are navigated. Christensen’s aim was not to assume the status of child but to walk a tightrope between being recognised as an ‘adult’ while avoiding association with any known category of adult, and the accompanying preconceived ideas, practices and connotations associated with that category, as well as adulthood more generally. For example, while following some set ‘rules’ for adults within the institution, Christensen would also follow children’s ‘rules’ and practices in participation with children—when joining or leaving a game—and avoid traditional adult roles such as correcting behaviour, resolving disputes or looking after children. As this ‘different sort of adult’, Christensen did not attempt to minimise typical adult ways of relating to children, instead seeking to understand their views and wishes and treating these respectfully and as important.

Anthropological perspectives on researcher roles

Writing from anthropology, Christensen brings a sense of ethics as an ongoing, relational and performative process; something that can be missing from childhood studies where ethical discussions tend to be grounded in universal definitions of childhood and children’s rights (Meloni, Vanthuyne, and Rousseau 2015). However, it is this anthropological view of how generational differences are codified into modes of relations in the field that usefully illuminates implications of the researcher role. For example, teasing in the Tūrama School context is a common and usually benign relational mode between adults, but from adult to child can function as an assertion of power or socialisation technique (Morton 1996), and between children represents a form of cruelty. Although teachers frequently teased children, the hurt looks when I tried gentle teasing (with children I already had good relationships with) told me that I needed to reflect more on how the implications of relational modes shifted with my positioning.
In this anthropological view of the ethical as relational, the role of the researcher becomes understood not as a predetermined, static set of rules that the researcher brings with them into the field, but as a plurality of relationships actively negotiated through every day mundane interactions, dynamic and shifting, and working both ways, as the role of the researcher influences participants’ modes of being, power relations, and ways of seeing themselves, and vice versa. The field is an ecology of dynamic relationships and interactions, which is altered without volition by the researcher’s presence (Dominguez 2016). These characteristics are not unique to research with children, but are foundational to unpacking the ways in which researchers play into or disrupt the generational power dynamics that are specific to child research. The researcher may reinforce or undermine the previous power structures, including teacher authority and also power dynamics within peer relations. The researcher brings new social capital that is disseminated in unequal ways and is used and manipulated by children and adults to reconfigure power dynamics. Other authors have noted the potential for children to take advantage of the researcher’s privileges, and this was my experience as well—for example when I was invited to play tennis, only to realise it was because the children had been told they were no longer allowed to borrow tennis balls, but I as an adult would be. At other times, it was the teachers who would use me to support their agenda, in strategies to shame or control students.

Thinking in relational terms also helps with integrating the ‘scientific and moral aspects of knowledge production’ which Meskel and Pels (2005:3) argue represents a more anthropological approach to ethics. In the postmodernist rethinking of epistemology as dialectical, knowledge itself is co-constructed through interactions; the research process is not so much exposing knowledge as making it (Clifford and Marcus 1986). This is a dynamic central to fieldwork, what John Borneman (2009) calls a ‘dialectical objectification’, whereby through encounters, interlocutors make each other as subjects and objects, coproducing unique knowledges from the subsequent insights. Because of existing dynamics of authority between adults and children, however, adults may start overwriting children’s knowledge; when I played back early interview recordings I noticed myself following up on what was important to me, and missing what else was said, jumping into silences loaded with meaning, and interpreting children’s utterances through my own adult lenses, without considering alternative ways of understanding. Furthermore, in a school context where I was there as a classroom helper, I was very aware that I was shaping children’s knowledge at the same time that I attempted to study it. A little way through the interview process I realised I was still
trying to hold onto a purist concept of research and the researcher, attempting to elicit children’s views without ‘tainting’ them with my own perspectives. Yet knowledge is inextricably woven into relationships, produced and used through relationships, functioning as a commodity, capital, or as a gift, and especially intrinsic to adult-child relationships where the control of knowledge works to maintain power dynamics (Foucault 1990). Slowly my interviews turned into dialogues with knowledge shared from both sides. The children seemed to value my explanations of antibiotic resistance and the knowledge that they had had strep throat, not rheumatic fever, and in turn their responses help clarify their knowledge for me. As much as I may try to represent what children know and how they know it, this cannot be parsed from how I came to know what they know, and how I contributed to what they know. Ethical research, therefore, involves a process of paying attention to the kinds of understandings that are produced collaboratively within varying researcher-participant relationships; embedded within, rather than supplementary to knowledge production (Pels 2005). The picture book is a material product of this process, the knowledge within it a synthesis of their emic and my etic perspectives, and of the relationships I had with these children—including the fact that I held the greatest social, as well as editorial power.

If ethical and scientific rigour are grounded in relationships, then this is where the kind of adult makes a difference. When Mayall was attending to, rather than minimising generational differences, she could focus on understanding how children work within the social order. When I took a more authoritative role—for example, when asked to be a leader of a group of children on a school trip, I experienced one side of childhood: the mode of relations with adult authority. I became the constraint they pushed against to run and shout and play, while I felt the dirty looks of museum staff. I learnt a lot about how children relate to this kind of adult that day, as well as how museums are not designed for children. Other times I could be a ‘guest’ child, joining in their games. When a group of girls invited me to participate in their lunch time game involving rounds of questions, in sharing my own answers, I learnt what the girls most feared, their favourite movies, and about their crushes on boys. My adultness came back with a mention of my partner, and I then learnt how they perceive the lives of women; they act out a proposal and shower me with grass confetti. Attempting to shed adultness, as in the least-adult role, therefore limits the knowledge produced, because it denies the realities of children’s relationship to adults, the adult privilege the researcher holds and how children might be using it to their advantage, and the adult agendas the researchers bring with them which shape the choices being made; who the researcher follows and what is noted down or
remembered. Perhaps counterintuitively, in accepting adultness, the ‘unusual adult’ role allows for greater attention to what kind(s) of adult the researcher embodies, using adult identity as an analytical lens for interpreting both researcher-participant relationships and the knowledge produced within them.

The unusual adult role is not without ethical problems, however. While all of the authors mentioned (particularly Mandell) acknowledge practical challenges in maintaining their roles within a field twisted with complex power dynamics and multi-dimensional practices, when I entered fieldwork at Tūrama School with the ‘different kind of adult’ role in mind two problems emerged. Firstly, setting aside conventional adult roles can become politically ‘sticky’ when researchers come across children’s taboo behaviours which would normally be policed by adults. Secondly, the ambiguity of a researcher role can mislead child participants. In the remainder of this chapter I detail how these two problems emerged to challenge my positioning in the field, and propose the concept of ‘transparent guest’ as an operational mode for negotiating such issues in fieldwork with children.

The unsanctioned activities of childhood

The first problem arises with the discrepancy between how the adults in the field expect children to be, and how children actually are. Authors have noted there is a darker side to children’s private worlds that Western adults are not comfortable with as they do not fit within the neat construction of childhood as an innocent and ‘nice’ world (James and Jenks, 1996; Christensen 1999). Accessing these areas of children’s worlds can become wrought with political, social and emotional tensions when researchers are seen by other adults as not acting as adults should—condoning or colluding with children’s ‘inappropriate’ behaviour by failing to intervene or disavow, or even participating in the behaviour.

Both Mandell (1988) and Christensen (1999) describe ethnographic moments of stumbling over children’s practices that were unsanctioned by adults. In the day care where Mandell conducted her research, urinating in the yard was a regular, but forbidden practice. Mandell’s solution, when children tried to involve her, was to leave the scene. Similarly, Christensen (1999) writes of her encounter with a group of ten-year-old boys during fieldwork at a Danish school who show her that they have taped up another boy, Danny, onto the floor of a cloakroom. She describes in detail the dynamics of the scenario; Danny struggling to free himself while smiling and laughing, the excited, but not aggressive demeanour of the other
boys, the contrast between her confusion and unease and the relaxed and unaffected responses of the boys involved, including Danny himself. And she describes her internal conflict as an anthropologist; to act as an adult ‘should’ and be responsible and protect children, to intervene, or tell another teacher, and lose the trust of her informants? To casually suggest another teacher might like to walk through the cloakroom? To ignore, and by default condone, the behaviour? Yet Christensen is also aware of the privilege she had been granted, to be ‘let into’ the secret world of boys, usually inaccessible to adults, and that here was also an opportunity for deeper understanding of children’s social worlds, but also a desire to reciprocate the trust that had been given with loyalty. Christensen asks the boy on the floor if he knows how to stop this if he wants to, and Danny says he does, and the other boys confirm that at that point they will let him go. She walks away; eventually the boys emerge from the cloakroom, apparently the best of friends again, their game is over.

This encounter ends up significantly shaping the direction of her research, and Christensen reflects on a number of ways in which her experience problematises notions of adult and child and ethical research. She notes that the event from the child’s perspective did not fit within adult dichotomies of ‘nice’ and ‘nasty’, and nor did it align with adult conceptions of children’s relationships with each other, and with adults. In the position Christensen found herself, the event shed light on cultural expectations of the responsible and protective adult and vulnerable children, while demonstrating how these adult perspectives are not necessarily shared by children. But of most relevance to my discussion here, the encounter problematised the nature of exchanges between researcher and informants and highlighted a conflict between children’s cultural values of peer trust, loyalty and comradeship, and adults’ cultural values of adult responsibility and protection. This is where child research becomes politically thorny: the reality that children do not conform to adult expectations of what childhood should be, together with the expectation that the responsible adult will police these instances where children slip out of what children should be. This adult role is what makes children’s secret worlds secret, and what makes these so challenging for the adult researcher to access.

My introduction to this secret world, and the political challenges that came with it, came at the point with the relationship shift from participant to friend—a transition which Dominguez (2016) notes already raises ethical questions in research with adults, let alone children. My

2 Both sets of cultural values are also embedded in the principles of ethical guidelines: confidentiality versus protection.
inclusion in the wider network was gradual and happened with some children before others. One day ten-year-old Pikau was asking me to sit next to her on the mat; a few days later another girl was, and then a socially ostracised boy, and a couple of weeks after that another group of girls were making room for me to sit down. So when one morning I arrive at morning tea and am invited to sit with a group of girls while they eat, I think nothing of it. I am curious about the particular combination of girls, since the group includes Pikau, who did not have strong friendships when I first arrived, and Eponi, who lingered at the periphery of groups, but also Ruby and Tiana, who I had always observed with the popular girls in the class. After the bell rings for play-time, they invited me to join them. ‘What are you doing?’ I ask. Just hanging around and talking and singing, they say. We lounge around the side of a building and sing some of the latest pop songs and then some waiata [Māori songs]. As the bell rings and we head back to the classroom, I ask Pikau why Tiana is playing with us. ‘She broke up with her friends’, comes the reply.

At lunch, I am invited to join the group again. Thinking it will be more of the same hanging out and singing, I come along. Instead, I am told we are playing ‘dares’. This is a familiar game from my own childhood, and I quickly pick up the rules of this version; we all simultaneously throw a ‘paper-scissors-rock’, and if only two of the three hand signals are thrown, then the victors get to ‘dare’ the losers. I am curious about the game and interested to see how the dares compare to when I was a child. It becomes apparent that the girls’ dares are mostly focussed on daring a girl to ‘ask out’ a boy—usually an unappealing boy—and if he says ‘yes’ the girl would have to ‘go out’ with him [be his girlfriend].

I am aware that the adults at the school would not view this as appropriate play. A few weeks prior, Mrs Randell had given the class a stern lecture about how she didn’t ‘want to be hearing any more of this boy-girl nonsense. You’re all far too young for that’. Yet I also know that these sorts of themes are an important part of children’s cultures—I remember them from my own childhood—a kind of ‘revolt of the sexual body’ (Foucault 1980:57) in response to societal suppression of sexuality where the disapproval of adults adds illicit associations to the practices, reinforcing them as symbols of in-group status. However, it is one thing for me to turn a blind eye to things I might see as an outside observer; it is another for me to be a participant in things that would be considered inappropriate from the perspective of adults. I find I am unsure about where the line is, and whether I can ‘semi’ participate or whether I need to withdraw from the game completely, and lose my in-group
status as a consequence. When I lose the paper-scissors-rock game, I say to the group, ‘hang on a second girls, I need to talk to you about something first.’

They gather around, and I say, ‘You know I’m here from the university to do research with you guys, right?’

‘About asthma,’ says Pikau.

‘Yup that’s right. So I have to be a bit careful about what I do because I could get into trouble with the university. So I can’t like, ask boys out or anything.’

This explanation is readily accepted by the girls, who decide my dare will be to ‘twerk’. I have no problem with making a fool of myself, and I go ahead and dance for them with exaggerated effect. They laugh uproariously. When it comes my turn to dare, I carefully keep my dares benign: dance around the courts saying ‘I am a beautiful ballerina, watch me twirl’ and go into the playground and sing the school song. These seem to be acceptable dares and get shouts of approval from the girls who are not on the receiving end, while the girls who have to carry them out blush and squirm and complete the dares in shame. I am curious about their behaviour which contrasts with the attitude I had taken to my own dare. The dares are mildly embarrassing but not hugely so, and most other students aren’t even paying attention.

Throughout this I am experiencing a mixture of feelings; unease about whether it is okay for me to participate in this way, delight that I am being included, and fascination with what function this game is serving and what meaning it has for the participants. I am conflicted by my desire to be accepted, my anthropological interest in the interaction, and my unease with playing a role other than the responsible adult.

This unease is exacerbated as we sit chatting on a picnic table and a teacher on duty comes by and asks us what we are doing.

‘Talking,’ one of the girls answers.

‘Well that’s not what you're supposed to be doing at play-time, is it?’ replies the teacher. I watch this interaction, baffled. Are they not allowed to talk at playtime?

‘But we’re not allowed to talk in class, so we have to talk now,’ explains one of the girls.

‘You need to talk while you’re playing, then,’ instructs the teacher. ‘Go and play tag or run some laps.’
I am silent, stiff and bewildered about these strange rules I am not aware of, and wondering what the teacher is thinking of me. We move off, as if to do as the teacher said, but then double back and walk into a new area of the playground to continue our game. I have lost the round of paper-scissors-rock and it is my turn to be dared.

‘You have to go and twerk in the bathroom and you can’t stop until I say so’ orders Ruby. I am taken aback by this dare—it is clearly calculated to push me much further than the last one I did. I am uncomfortable with going into the bathroom, I am not sure if I am allowed to be there as an adult. I have not had the rules explained to me. Most adults would not find themselves in this situation. Ruby laughs at my uncertainty and pushes me into the bathroom. I suddenly realise the importance of shame—and the performance of shame—as central to this game. I had not displayed a sufficient level of shame the last time I had done the twerking dare, and as a result the girls were significantly upping the ante. I quickly performed a half-hearted dance and escaped the bathroom, proclaiming my embarrassment. Despite the complaints from Ruby that she ‘hadn’t said stop yet’ my performance was met with approval. Although I didn’t complete the dare, the actions have met their purpose; to shame me and as such, demonstrate my belonging to the group.

The next child dares Ruby to dance around a pole like a stripper. She does so with a finesse clearly learnt from music videos. My adult self is shocked; my child self wants to try it too. The other girls show off their pole dancing moves and the bell rings.

After school finishes, I drive home uneasy about my level of participation, but also fascinated by what I had witnessed. What was that? By the time I get home I have a theory; the emphasis on shame and performance suggests I have got myself caught up in a kind of initiation process for a newly forming group.

The next day, after a prize-giving assembly the deputy principal adds an announcement reminding the children they should be playing a game or doing something active at lunchtime. I realise this is where the duty teacher yesterday was coming from, and that I had been inadvertently undermining the school’s attempt to regulate the children’s free time. Yet knowing this put me in an awkward position when at lunch I am again invited to play with Pikau, Ruby, and Tiana, and they clearly have no intention of complying with instructions. I go along with them again, but I am nervous of a duty teacher seeing me and thinking I was condoning or encouraging their inappropriate play.
Today Eponi is not with us. I ask the others where she is. ‘We’re not friends with her anymore,’ came the reply. ‘She didn’t do her dares and she complained, so we kicked her out of the group’. This must have occurred earlier at morning tea. The absence of Eponi tells me that I might be right about my thinking that this dare game is a kind of initiation process. The game starts again immediately after we finish eating, and I make a performance of reluctantly letting them laugh at me twerking again (this time I decline to go into the bathroom) and suggest a few dares of my own when it’s my turn: ‘do a handstand on the grass.’ One of the dares is to ‘flap your arms and say ‘I am a retard.’’ ‘Aw that's mean' I say, but don't comment further on the language.

The game seems to run its course, and, having served its purpose of establishing us all as group members, is retired. To my relief, we only hang around a picnic table and sing waiata for the rest of playtime. I am able to Google the words to songs on my phone and am conscious of the social capital I hold as an adult who is allowed to carry a cellphone.

The unease that lingered after the bell rings reflects the disjuncture between the way adults expect children to be, and the way children actually are, and expectations about the way
adults are supposed to relate to children. I felt uncomfortable being witness to children’s games that did not match the adult idea of what is appropriate play, including play that explored themes of shame, power and sexuality, and uncomfortable that I was observing and participating rather than discouraging or policing. This was the first of many occasions when I had to navigate such political awkwardness. Yet I had learnt more about these girls in two lunchtimes than I had in the preceding weeks of peripheral observation. Although at Tūrama School there was an especially stark difference between how adults wanted children to behave, and how they actually are, in any fieldwork with children the adult researcher role has potential to create problems with consequences greater than just discomfort. The times and places where this issue manifests may usefully highlight adult cultural conceptions of childhood in different contexts, but the challenge is to navigate these moments without giving primacy to these adult conceptions, and while maintaining good relationships with both adults and children.

The transparent guest

The experience of this event shaped my approach to the remainder of my fieldwork through the discovery that transparency about the rules of my role was an effective way of placing boundaries where I needed to. When faced with the possibility of dares that could put me in a compromised position, simply explaining my position to the girls gave me a legitimate explanation for non-participation. While on the first day I was caught by surprise and let myself be pushed into the bathroom, on the second day I was able to explain that I was not allowed in the bathroom, and that I might get in trouble with my university and maybe wouldn’t be allowed to come to their school anymore. In this, I took on a new role: ‘transparent guest’, openly acknowledging my outsider status, and using the norms that the children themselves understood to negotiate my position and justify my actions. With the guest role comes an established set of shared rules that children themselves understand well. Guests, ideally invited, enter and observe in order to learn and follow the rules and cultural practices of the host. But guests may also be exempt from some house rules, or understood as naïve to host rules and practices and so transgressions are more easily forgiven, though if some rules are broken the guest may not be invited back. Guests may also be bound by their own rules brought from home, and in children’s culture, ‘my family say I’m not allowed’ is a valid excuse for non-participation. As a guest, I could move in and out of different contexts, adapting to the new norms and rules as appropriate. I was sometimes guest adult, but when
invited, could also be guest child. Of course, guests are also seen as special, at least initially, but over time and frequency of visits can become mundane, ‘like family’, and see more ‘realness’ from the hosts.

As a principle, ‘being open and honest’ is generally applied in terms of transparency around research: aims, funding, implications (Price 2016). Here I extend transparency into relationships: as much as possible making the rules which govern my behaviour visible so that they are understood and predictable, and in doing so, respecting my participants’ competency. I also took this approach with adults in the field. While other authors note struggles with adult gatekeepers and authority figures, few seem to address these directly, as if the researcher’s approaches must be kept secret. Shortly after this first event, I attended a staff meeting and took an opportunity to explain my role in the school in more detail, including the explanation that I might sometimes do things that look strange or unusual, because was trying to get alongside the children as much as possible. I also invited the staff to approach me directly if there were any questions or concerns about what I was doing. After this, my discomfort about playing with the children was much reduced, and sometimes talking through my ethical quandaries with teachers (for example, how I should respond when children begged food off me) seemed to help legitimise my behaviour as well. As a model for research with children, the ‘transparent guest’ role positions ethnographers as actively negotiating and renegotiating complex rules, including those that implicitly and explicitly govern the researcher, and in the most visible way possible in order to minimise ambiguities and forestall many of the tensions that can arise from perceived deviancy in practices.

Harmful implications

Transparent acknowledgement with participants of the rules researchers operate under is also important when considering a second problem arising from an ambiguous researcher role. Existing categories of adult identity come with known and predictable sets of rules that are politically important for assisting children to navigate the adult world. In separating themselves from these categories, researchers enter unchartered territory not only for themselves, but also for their participants who are learning how to interpret and respond to this new kind of adult. As others have noted, researchers demonstrate their role to child participants through their actions, and like others, I recited the mantra ‘I am not a teacher, go
tell a teacher’ until children learnt that I would not intervene in their disputes, or tell off rule-breakers. Another significant way that I expressed my ‘different’ adult role was by transgressing a simple norm and sitting on the ground with children, rather than in chairs with adults. In this separated world, my behaviour was not disapproved of—at least not overtly—but certainly noticed. When I sat with the class on the courts to watch an athletics competition, I was asked numerous times if I would like a child to be sent to fetch me a chair. When I sat on the floor of the school hall with the classes waiting for the bus to arrive to take us on a field trip, teachers and parents all suggested I might like to sit in a chair. ‘I like sitting on the floor’ became my stock explanation, and before long the children were explaining to adults ‘she likes to sit on the floor’. I was seen as odd and unusual, but also there was an understanding that as an anthropologist, it was acceptable for me to do strange things. From the children’s point of view, the simple act of sitting like them seemed to signal that they could act in their normal ways.

However, because these ‘differences’ are constructed, negotiated and learnt through interaction in situ, not everyone in the setting—including the researcher—holds a consistent, complete understanding of the researcher role. In adopting a role outside of social norms researchers inevitably will encounter misreadings, misunderstandings and tensions from both adults and children. Discussion of such problems in the literature has done little more than acknowledge these, focussing on how they might cause discomfort for the researcher who receives funny looks or jokes from adults, or loss of positive relationship, for example, when researchers have declined or avoided helping adults in the field (e.g. Mandell 1988). Thorne (1993) worries that by participating—or even simply declining to intervene—in children’s rule breaking she is undermining teacher authority and the social order of the classroom.

Less acknowledged is the potential for real harm caused by the ambiguity of the researcher role to others in the field. Mandell describes an incident where she became so lost in the least adult role that she neglected to intervene when a child cut open another child’s head with a spade. Mandell focusses on her embarrassment at the scolding she receives from the teacher on duty, and how quickly she was forgiven. She does not question whether her presence as an adult meant that the teacher did not step in as soon as she might have had she seen only a group of pre-schoolers, or if other adults patrolled the area she was in less closely. She also didn’t comment on what her prompt pardoning indicated about her privileged position in the day care, or who ultimately bore responsibility for the injury.
The problem with setting aside the normal rules of known categories of adults is that the limits of this role are not made clear until they are reached in the moments of enactment. While visibly forgoing some of the usual rules of adults, I would still abide by others. From the perspective of the children, this must have seemed quite arbitrary, and may have made me difficult to predict. Where this became most apparent was when I became privy to talk or behaviour that children would routinely keep from adults in order to avoid adult intervention; notably instances of abuse (‘hidings’), suicide ideation or self-harm. On the one hand, this is the aim of child research—to get at those hidden practices and develop a child-centred understanding of their world. On the other hand, children keep those worlds hidden for a reason, while I am ethically bound to reveal them. The problem was not deciding whether or not to break confidentiality in each instance, although this was difficult, but in realising that I may only be coming into this information in the first place because I had set myself up as someone who did not follow normal adult rules. I was in essence misleading children about how they could expect me to respond in other situations.

This problem erupted into my awareness when during an interview with two boys, nine-year-old Whetu cheerfully described how he didn’t care when he got hidings from his mother with a pole, with a belt ‘with the metal bit on top’, and with a broom ‘and the broom broke’. Relaxed and chatty, he seemed unaware of the implications of his disclosure for me; when I asked if anyone else knew, he listed a group of friends, ‘just people that won’t tell’.

Physical punishment of children has been illegal in New Zealand since 2007 (Crimes (Substituted Section 59) Amendment Act 2007). Most children are well aware of this, and also infer that if their parents are reported, they may be jailed and the children removed from their care. As practices of physical force still persist in many homes, this puts many children in the awkward position of protecting their parents, even while suffering. The children of Tūrama School are very familiar with CYF, the agency responsible for the protection of young persons under 17 years old. On one occasion in the classroom I overhead one child correct another on the proper preposition: ‘you don’t go to CYFs, you’re with CYFs.’ This linguistic familiarity reflected the high level of contact CYF had with children at Tūrama School—CYF workers also frequently visited the school as part of interventions with families. CYF is less familiar to children in less challenged circumstances—the agency was never mentioned in interviews with similarly aged middle-class children during my master’s research (Spray 2012, Spray et al. 2018), for example.
The point is that by eight or nine years of age Tūrama School children are well versed in the consequences of letting information slip to adults in the school, and either have themselves been (or are currently) under CYF care, or have heard from peers about the consequences of CYF involvement. As well as fears of being ‘taken away’ to live with strangers, children fear getting their parents in trouble, and retribution or shame for betraying their family. As a result, children tended to be circumspect, and though talk of ‘hidings’ was ubiquitous, it was always generalised, or talk about someone else. The school social worker, Lucky, would liaise with families who consented, and reports to CYF were made when necessary to uphold legal obligations.

Unbeknownst to Whetu, I was bound by my university ethics agreement, as well as human responsibility to act on the information. Despite my obligations I agonised over what felt like a betrayal of this child’s trust, of breaking the ‘children’s code’. I did not take Whetu’s bravado at face value; if his intention was to shock and impress, there were also indications of distress underneath the smiles: ‘but it’s crying’ he said, to laughter from his friend Jackson. He cries, but his options constrained, he copes with talk about how little he cares. But, in the moment, with his filter off, he did not make a conscious decision to disclose that information, in full knowledge that in my role as adult, it would be passed on. I had spent the whole year demonstrating how I was not like other adults; I would tolerate or even laugh at their swearing and dirty jokes, eat their forbidden lollies, shrug off talk of mischief, gangs, and stealing, and turn a blind eye to graffiti or vandalism. Although my participant information sheets said I would break confidentiality for safety concerns, why would a nine-year-old remember that after months of experiencing me in this different kind of adult role? I felt I had misled him, and now if his situation worsened it would be a direct result of my naivety.

If this were an isolated incident, it might be easier to think Whetu did want me to know, and did understand the consequences, or that this was a rare and unfortunate side effect, perhaps even in the child’s best interests. Although this was the most serious, there were a number of other occasions where children disclosed information to me that I am confident they would have been more cautious about revealing to another adult. Later in the year a number of girls showed me self-harm marks and then, upon seeing my adult-like response, begged me not to tell a teacher (I told). I also flagged concerns after another disclosure of ‘hidings’ by another boy’s father.
In the case of Whetu, I decided that since my lack of transparency had led to the disclosure, I should at least be transparent about what I was going to do about it—even if it cost me rapport with him and probably his friends. I asked to pull him outside of class one day, and while he wound himself around the veranda post and avoided eye contact, I explained that I was concerned about what he had told me, that it sounded like mum might need some more help, and how would he feel about talking to Lucky about what was going on. Still looking everywhere except at me, Whetu smiled and said Lucky was cool. ‘Can I go now?’ He made a rapid escape, and I figured he’d never speak to me again. I did not learn where the matter went after I gave the details to Whetu’s class teacher to be handled privately through school channels. I was surprised though, when a few weeks later Whetu and his friend Leo (who had until now largely ignored me, I assume because he had been involved in misbehaviour) came running to meet me before school, dragging me around to visit friends, classrooms, and play basketball. My popularity with the pair continued for the rest of the year. It appeared I had earnt rapport, rather than lost it in this instance. There can be an ambivalence in children’s relationships to authority; the potential for options children cannot access on their own, but at the expense of control and carrying the fear of over or under response. It seemed like my actions with Whetu had situated me in the ‘right’ kind of adult category. For these boys, caught between craving and fearing adult attention, my interventions may have ended up being more helpful than harmful, though I do not really know, and a positive outcome in this situation does not preclude the potential for harm next time. The problem I am concerned with is the degree to which I found myself in a position of my own making, taking away children’s agency and autonomy when they were likely not aware of the consequences of their disclosure. The nature of the issues children at Tūrama School deal with makes this a higher-stakes problem than what might be encountered at a middle-class school, but similar dilemmas could be created in research any place where exist great discrepancies in power between adults and children.

Conclusion

Dilemmas and tensions stemming from generational differences are commonly noted by child researchers, in passing comments, or cataloguing of challenging experiences. Yet these have not tended to be the subject of analysis, unpacking questions about why these tensions emerge, what they tell us about childhood and adulthood, and what role the researcher plays in their creation. Nor is there much discussion of what researchers should do, how conflicting
responsibilities may be weighed up, and the implications of various responses. How does the way the researcher role is negotiated create new ethical tensions? Whose concept of childhood are researchers using in decision-making, and is this because it is the most ethical, or simply belongs to the most powerful? When researchers choose to walk away from rule breaking or taboo play, rather than observe, participate, or police, what impact does this have on their relationships with their child participants and the adults working with those children?

My intention in this chapter was to go beyond acknowledging difficult circumstances in fieldwork by clarifying how and why these difficulties arose, considering the nature of adult-child relationships in this context, how childhood and adulthood are conceptualised, and how children perceive and engage with adult roles. In doing so, my aim is not to solve these issues—I do not think they can be—but demonstrate how thinking about generational issues in fieldwork through a relational framework can help researchers better understand the implications of their role, and find small strategies—like transparency—to mitigate these difficulties.

Having established the context for this study, and the methodological processes and challenging the collection of data, I now turn to an analysis of children’s health coproduction at Tūrama School and surrounding community. The subject of child and adult identities continues in chapter four, where they manifest in authority structures and relations of care within the school which shape children’s practices of the body in patterned ways. The nuanced understanding of child and adult relations I gleaned from having to work within them has therefore also come to bear significance to understanding the dynamics of children’s health.
In this chapter I begin to answer the main question of this thesis: how do children participate in the coproduction of their health? As discussed in chapter one, the anthropology and sociology of childhood literatures have long established that children produce cultural understandings and practices in relation to health and the body (Bluebond-Langner 1978; Christensen 1999; Clark 2003; Mayall 1993, 1996; Prout 1986; Prout and Christensen 1996). This literature has not always captured the processes through which these understandings and practices are guided or constrained by the structures of society, nor the ways in which these structured practices come to shape children’s health in patterned ways. In this chapter I build on this literature by elucidating some of the processes through which children’s meaning-making and practices articulate with, and are structured by the institution and state health interventions. I centre on the Tūrama School clinic for rheumatic fever prevention and related media campaigns as an entry point for addressing how children engage in interpretive practices in relation to adult-child relations, the body, the school institution, and public health policy, and how these practices, in turn, can shape children’s health.

Fundamental to answering the questions of this thesis is the context of children as peers within an institution marked, as discussed in chapter three, by distinct power structures separating child and adult roles, rules, and authority. Importantly, schools are also the site at which the state has arguably the most direct relationship with children, where the curriculum and daily activities are to a large degree guided by state policy, and where interventions targeting ‘at risk’ children are strategically implemented to maximise efficiency of reach (Burrows and Wright 2007). The story of how children come to creatively appropriate a clinic into their social practices and produce new understandings of illness is therefore also a story of how adult-child relations in the classroom can influence the implementation of state health services, and of how a public health policy can restructure the processes through which children learn how to experience their bodies. These are complex and multi-faceted dynamics, and so in this chapter I touch on themes that will be developed over subsequent chapters: the way children’s strategies to stabilise their social position may conflict with adult agendas and perceptions of what child wellbeing means; the way children’s activities bring
state policies to life in the field and can powerfully impact their effectiveness; and the way that modes of service delivery can restructure children’s interpretations of their body.

In this chapter I focus on two main processes through which children’s health is coproduced. Firstly, I elaborate Christensen’s (1999) analysis of how children collectively construct illness knowledge by demonstrating how state interventions can restructure this embodied process in such a way that children produce novel understandings of ‘sore throat’ and ‘rheumatic fever’. Secondly, I explore how, structured by the institutional context of the school, children collectively appropriate the clinic into the formulation of social routines which mediate their engagement with health care. I identify three motivations for these routines, none of which necessarily represent conscious, nor mutually exclusive strategies. First, as navigating peer relations is a salient concern of children (see chapter seven), children can appropriate the clinic into strategies of relationship-building. Second, children’s agendas can often conflict with those of adults, creating tension between authority structures and resistance that can result in stressful classroom environments, so that children may use the clinic as a place of respite. Third, by creating social routines of ‘getting out of class’, children may also be creating new ways to legitimise access to health care while maintaining ‘tough’ reputations among peers. Both children’s creative conceptions of illness and the social routines they construct around the clinic mediate the ways they access health care from the clinic and, on occasion, their use of pharmaceuticals and other health practices—key ways in which children contribute to the coproduction of their health.

“Who has a sore throat?”

It is the end of summer in my early days of fieldwork, and children entering the classroom after the morning bell are breathless and sticky from an early game on the field before school. Mrs Randall’s classroom is brightly decorated with student artwork, carefully mounted on black card, labelled with the neat, round handwriting of teachers, and pinned onto the wall or pegged onto strings that bisect the airspace above us. As we do every morning, we sit on the mat, a large carpet on an open area of the floor. I sit towards the back so my looming body does not block the view, and children slip snugly on either side of me even though there is lots of room, leaning against me, their bodies damp and hot. As Mrs Randall takes the roll, Jackson and Aaron are pinching the sides of the boys sitting in front of them. The boys flinch
and try not to squeal, keeping their faces trained to the front so as not to draw their teacher’s attention.

‘Who needs to order a spare lunch,’ Mrs Randall asks, and writes down the names of two children on a paper form. ‘And who has a sore throat?’ Over a third of the hands go up. I glance at her to see if this is normal, but she is already writing down the names.

‘Whetu, take this to the office please darling,’ Mrs Randall says, handing him the forms. A couple of latecomers slip into the classroom: Verity and her younger sister Karina. Mrs Randall’s eyes widened. ‘Why are you two late?’ she exclaims. Karina looks embarrassed but Verity shrugs. ‘Our mother was late,’ she says. Mrs Randall looks like she might say something further, but at that moment another latecomer appears at the door. Mrs Randall appears to reassess the battle and only sighs.

‘Alright. We are going to work on our sample books now,’ she says. Several children, including Verity, bounce up and head for their desks. ‘Did I tell you to go?’ Mrs Randall barks, and they pivot, Verity dragging her feet and looking at the ceiling. Everyone sits down again.

‘Now you may go-’ and there is a mass break for the desks, overwhelming Mrs Randall’s shouts for them to go quietly and stop running!

The desks are organised in mixed-gender groups, facing each other to form a table, and I take an empty desk next to Caleb, who has started writing in his sample book using a thick, intoxicating black marker pen. Several times a term, the students complete a piece of work to be sent home to the parents, showcasing their learning, and Mrs Randall has given me a stack of the sample books to mark for spelling. Jackson, nearby, is swinging on his chair and making electric guitar sounds.

‘Be quiet!’ Mrs Randall snaps at the class, and there is a moment of silence before a dull hum resurfaces. Their sample book worksheets for term one are called ‘My School’ and the children have to copy out the school expectations and answer questions about themselves: things they have achieved; how they can demonstrate responsibility; ways they could be fair or not fair; who their role models are. The children at my table hold pencils over their workbooks and chat about their families. ‘My mum has too many kids,’ says Cassidee. I tune in and out of their conversation. Her mother is 45 years old. In the book I am marking, Whetu has supplied an example of when he has been fair: ‘My sister drew on my face when I was asleep so I drew on her face when she was asleep.’
‘Jackson, be quiet!’ Mrs Randall shouts, and the electric guitar noises pause, then resume.

At the morning tea break, Cassidee sits by me. She has a large sore on her finger: an inflamed, pus-filled bump. I say, ‘Ooh, what’s that?’ because it looks painful. Cassidee shrugs and moves her hand out of view, and continues eating her chips. Her classmates are talking about chilli. ‘I ate some one time and it was so hot!’

‘My sister woke me up once, because I was sleeping with my mouth open—’

‘-I drank some water but it didn’t help—’

‘-because I had a blocked nose—’

‘-and I was sweating—’

‘-and my sister put chilli in my mouth and I woke up because it was burning!’ We all laugh.

Back in the classroom, we do our spelling and times tables. I do the tests with them, but I don’t have to read out my result for Mrs Randall to record in her big book like everybody else. ‘Mason- Jackson be quiet!’ Mrs Randall barks, and Jackson stops banging his heels on the floor, and starts humming instead. ‘Nine,’ says Mason.

‘What?’ says Mrs Randall.

‘Nine,’ Mason repeats.

‘You only got nine out of twenty?’

‘Yes,’ says Mason. His ears are pink.

‘How can you only get nine? These are the ten and eleven times tables. The easiest ones! Jackson be quiet!’

‘The sore throat lady is here!’ someone calls out, and Mrs Randall throws up her hands and asks, ‘who do you want?’ Whaea1 Allison, the ‘sore throat lady’, reads out the names, and the children disappear out the door. I can see them through the window, running across the courts.

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1 ‘Whaea’ translates to ‘mother’ or ‘aunty’ and is used as an honorific indicating relative status of the speaker to the addressed; a close equivalent to ‘Mrs’ or ‘Ms’. The children addressed some teachers and staff with ‘Whaea’ or the male equivalent ‘Matua’, and others with the English honorific, depending on the person’s preference.
Rheumatic fever and the Tūrama School clinic

Rheumatic fever is a big problem out here, the principal had said to me in our first meeting before I started fieldwork. It was late 2014, and the clinic had been newly installed in a disused classroom, part of the government’s 2011 Rheumatic Fever Prevention Programme (RFPP), described in chapter two, to address rising rates of the disease among low-income Māori and Pasifika children. The clinic’s role is to swab children who present with sore throats in order to test for strep throat, the most common form of the GAS infection that can lead to rheumatic fever (Lennon et al. 2014). Children diagnosed with strep throat are prescribed a course of antibiotics, usually delivered by parents at home, with a sticker chart which functions both to incentivise children to take the medicine and assist families and the clinic staff to track its administration. Occasionally the nurses at school will administer children’s antibiotics when there are issues with adherence or when families do not have a refrigerator in which to store the liquid.

Around half the children turned up positive results for strep throat in that initial sweep, the principal told me—a statistic other teachers echoed. By the time I joined Mrs Randall’s class in 2015 the clinic processes had become integrated into the routine of the day. ‘You should be looking at rheumatic fever,’ one of the teachers advised, and within a few weeks of fieldwork, I had gained consent from the clinic staff and their manager to observe the clinic. On days when I was based in the classroom, I would follow the children with sore throats to the clinic; on other days I was based in the clinic and would act as runner for Whaea Allison, taking her list to each class and following the stampede of children back. The clinic staff also kindly offered me use of the clinic space to conduct some of my interviews, since the room was empty after school and quieter and more private than regular classrooms.

Not all schools in New Zealand have a sore throat clinic. Most school clinics are located in the Manukau or Northland regions, where rheumatic fever rates are highest in Aotearoa. The Tūrama School clinic is one of 61 operating out of schools across Manukau, targeting children deemed demographically ‘at-risk’—Māori or Pacific children in the lowest deciles of socio-economic deprivation (Anderson et al. 2016). The school swabbing programme operates in partnership with local providers, and with the consent of parents, approximately 95% of Tūrama School children were enrolled in the throat check programme at the time of my research. Whaea Allison, the whānau support worker or ‘sore throat lady,’ as the children refer to her, swabs the children to test for strep throat, while Deb, a registered nurse, will
attend to the positive results: liaising with parents, supplying prescriptions, visiting homes to drop off medicine if the parents cannot get to the pharmacy, providing information and support. In order to maximise swabbing efforts, the clinic has a two-pronged process. Every day, teachers ask their class if they have a sore throat as part of a suite of routine administrative activities, while twice a term, Allison will check the throats of all children in each class with a flashlight and send those with red or swollen throats to the clinic for a swab. Some of the teachers do not always remember to ask their class about sore throats every morning, and at times Allison has spoken with teachers to let them know their class came back with a high number of positive results in the class check, and to remind them of the importance of asking for self-identified sore throats. Mrs Randall, however, is consistent and structured with her routine; she takes her role in the fight against sore throats seriously, and just as well, because sometimes up to half of the children put up their hands to be swabbed.

While the clinical and political implications of rheumatic fever policy are to greater and lesser degrees the subject of policy attention, what is missing from official accounts of rheumatic fever distribution and subsequent health is the experiences, perspectives and activities of children themselves. Indeed, while the design, implementation and ongoing evaluations of the RFPP components have involved parents and caregiver perspectives, children fade into the background of these documents (Anderson et al. 2016; Ministry of Health 2013; Ministry of Health 2015; Vermillion Peirce, Akroyd, and Tafuna 2015), the passive sufferers of disease and recipients of care. As a result, the programme reflects parent understandings, concerns and barriers to care, but children’s experiences and perspectives are largely invisible. For example, evaluations recognise that parents do not typically think of children’s sore throats as requiring a visit to a doctor, and that standard practice is for parents only to ‘keep an eye on’ their child (Ministry of Health 2013). A large component of the promotional campaign has therefore been to emphasise that sore throats can kill. ‘It’s not cool to take the risk,’ goes the poster headline. ‘Get your child’s sore throat checked every time.’ The sad eyed child sitting in a hospital surgery on the poster is meant to incite parents to take their child’s sore throat seriously, but while they consulted with Māori and Pacific health leaders and tested the campaign on their target audience of parents, the Health Promotion Agency did not consider how children would view these images (personal communication). Other evaluations include the views of adolescents but not younger children, or only test children’s health literacy via survey (noting, for example, whether they have heard of rheumatic fever) while soliciting the full experiences
of adult whānau members through focus groups (Anderson et al. 2016; Ministry of Health 2015).

In rendering children’s experiences invisible, what policy and evaluation documents do not capture is the way the sore throat clinic inserts itself into the structure of the day, bringing with it specific notions of roles and responsibility for health care, and the constant, reinforced reminders to be aware of the throat. Policy does not anticipate the way that children will appropriate the clinic space into social routines of escaping the classroom, nor how public campaigns will contribute to processes of children’s health knowledge production and their understanding of risk and risk prevention. Policy does not consider how the daily practices of throat checking may reshape children’s experience of their body by teaching them to be hyper-conscious of the visceralities of the throat (detailed further in chapter six).

Likewise, apart from a small component of the RFPP which involves referring some families for housing support (primarily insulation), these studies of socio-economic disadvantage and disease neglect the well-established impact of psychosocial and homeostatic stress on children’s immune function (Haggerty 1986; McEwen 1998), including decades old evidence that stress may increase the likelihood of strep infection (Meyer and Haggerty 1962). Yet a child-centred perspective can immediately identify how not only structural conditions, but children’s interpretive practices, too, mediate the experience of stress in the body, whether through the psychosocial stress generated from peer relations (Spray et al. 2018, see also chapter seven), the way institutions structure bodily signals of cold or hunger (chapter six), or the way children’s social practices contribute to patterns of nutrition (chapter five).

All of these dynamics are embodied through a new habitus of children’s co-creation—the social structures and bodily dispositions that in turn guide their practice. In this way, an environment structured to disadvantage this group of children becomes physically embodied via the institution, while children's own practices, based on their interpretive understandings of that embodied experience, work towards generating new structures that guide the coproduction of health. A major government health initiative is connected to Mrs Randall’s classroom environment via the activities of children; activities which also contribute to the social structures that shape their engagement with health care and practices of the body. In the remainder of this chapter I discuss two of the processes through which children help to co-construct this new habitus: children’s embodied knowledge production and children’s social routines.
Visiting the clinic

The days are growing crisp and cool and with the temperature drop comes a settling into the school routine and an uptick of visitors to the clinic. Not all of the children who say they have a sore throat will be fetched from their classroom for swabbing today, though. With 20 classes, it was not usually possible to get around everyone, and Allison looks over the names with a discerning eye. ‘They’ve just had class check this week,’ she says of one class. ‘But maybe him—I don’t think he was there.’ She checks the names recorded in an exercise book. ‘These three I swabbed last week; they were all negative.’ This leaves Allison with a list to spend the day working through; she usually gets to about 20 students per day, but because I am here today to help with the running around, she may get through more. She hands me her list for Mrs Randall’s class, with eight or so names, and I make the long walk from the junior school, where the clinic is located, to the block of classrooms on the far side of the courts, past the hall, by the fields. I am greeted at the door with a chorus of ‘hi Julie’s. ‘Why are you here?’ asks Mason.

‘Quiet!’ snaps Mrs Randall. The Education Review Office is inspecting the school next week, and Mrs Randall has been in a frenzy getting her classroom in order. I read out the names of the children, who take off out the door.

The clinic classroom is set up with three low tables at the back and a bookshelf of books and a few toys to occupy children who are waiting for their turn to be checked. For some children, this is a space to be free, muck around, and talk unencumbered by the policing of teachers. If they get too rowdy Allison will reprimand them, but her attention is on the child she is working with and for the most part the waiting children are left alone until they are called for their turn.

I follow the stragglers into the clinic, where Whaea Allison is finishing up with the last group. The sprinters from Mrs Randall’s class are playing on the ‘spinning’ office chair in the corner, but nine-year-old Dandre heads straight for Whaea Allison’s desk, calling out ‘I’m first!’ Five or six others pull over chairs in a semi-circle alongside Allison’s desk, chatting and joking with her as they watch each other undergo examination. Under the mournful gaze of the two boys in the rheumatic fever poster warning of deaths that started from a sore throat, Dandre weighs himself, helped by other children, and calls out the number to Allison to write down. ‘Do you have a sore throat?’ Allison asks, and he nods, opening his mouth wide for her to check with a small flashlight, and watching as she circles ‘sore throat’ and
‘redness’ on his form. He sits upright, beaming as Allison lets him check his own temperature with the electronic thermometer, and holds statue-still while she swabs the back of his throat—a process all children describe as uncomfortable since it ‘tickles’ and can stimulate gag reflexes. ‘Well done Dandre,’ says Allison; she has a friendly rapport with these children and Dandre basks under her attention. ‘Can I go back to class now?’ he asks, and waves goodbye to his classmates as he slips out the door.

When the rest of Mrs Randall’s class have gone, Allison sighs. Earlier, she and Deb had discussed a current dilemma—a large number of children who self-identify as having a sore throat on almost a weekly basis, but rarely test positive for strep. The problem varies by class, and Mrs Randall’s class, along with a couple of the other senior classes, has one of the largest
groups of regular clinic visitors. The main issue is the expense; each swab costs about NZ$15 and Allison feels an obligation to manage government resources responsibly. On the other hand, she doesn’t want to dissuade children from coming if they genuinely have a sore throat—the risk of rheumatic fever is real. Yet Allison’s suspicion is that many of these children were just here to ‘muck around’ and get out of class. ‘They go straight down the back going ‘I’m last’—they’re not really here to get their throats checked,’ she grumbles, although this was not the case with most of Mrs Randall’s class today.

This observation is supported by district-wide evidence which shows that while swabbing numbers had increased over the period, the proportion of swabs returning a positive result has decreased (Anderson et al. 2016). It has been noted in the literature that in regular clinics patients will fall within a normal range from avoidant to over-vigilant, and studies of school clinics indicate the same pattern. The term ‘worried well’ describes the 5–10% of patients who seek health care but who are not unwell (Korbin and Zahorik 1985; Lewis et al. 1977). By comparison, over the first seven months of my fieldwork around the same percentage of swabs for self-identified children—9% of 1434 swabs (range: 2–20%)—came back positive for streptococcus A (data provided by Manakidz). This does not mean that ninety percent of the time children are well—there are many other causes of sore throat—but these numbers do correlate with Allison’s and my own observations of children in the classroom. While I could not calculate the distribution of swabbing across children at Tūrama School due to privacy issues, the raw number of swabs the clinic took over the course of my fieldwork—3159 from both self-identifications and class checks—works out to average seven swabs a year for each child. In other words: children come to the clinic a lot.

A few days later, Dandre approaches me and asks why I didn’t get him today. Mrs Randall wrote his name down this morning, he says. ‘Because Whaea Allison said you already had a swab come back positive, so she’s going to get you some medicine,’ I answer.

Dandre looks confused. ‘But I have a sore throat today.’

‘Mm yes, but did you have a swab a few days ago?’

He looks uncertain. ‘Yeah I had a sore throat a few days ago.’

‘Yeah, so Whaea Deb is going to give your mum some medicine for you.’

‘But why don’t I need to get checked today?’
Because you already got checked, and you’re going to get some medicine soon.’

‘Julie, come play with us!’ Dandre is still frowning as I am pulled away.

Dandre’s anxiety on this day is what first prompted me to really wonder about the nature of children’s engagement with the clinic. It did not seem as if Dandre just wanted to get out of class, although I could also see what the nurses meant about the children who played and joked and tried to extend their time in the clinic as long as possible. It was not until I began interviewing children later in the year, however, that I could start to interpret how their practices of attending the clinic might be shaping, and be shaped by, their production of illness knowledge.

Learning sore throat

Working from an adult model of illness, a basic assumption of the RFPP is that children share an adult understanding of sore throat, and if children have a sore throat, they will know. This is an assumption I initially made myself, and I had some very confused moments in interviews before thinking to ask ten-year-old Anton, ‘What is a sore throat?’

‘Like the fever, you know? Coughs and that, sneezing and that. Cold and that.’

After I started asking, I got all sorts of descriptions, some of which represented a traditional or clinical idea of sore throat, but many of which did not. Alzea (age 10) thought the flu and sore throat were the same, associating both conditions with coughing, headache, and blocked or painful swallowing from coughing too much. For Trystan (11) a sore throat was ‘when you’re coughing and stuff. And your throat gets hot and that.’ Navahn (11) saw it as an allergy where ‘you can’t eat’ and ‘there’s lumps’. Meanwhile, children who had connected a painful throat to the label ‘sore throat’ often did not differentiate between sore throats with known causes, such as ‘talking too much’ or ‘eating scratchy food’ from sore throats that could represent a symptom of illness.

The processes through which such understandings of sore throat are produced might be best understood through a conceptual frame that places the experience of the body central to the production of illness knowledge, but also incorporates how the body comes to be interpreted through interactions with others. In her work with Danish schoolchildren, Christensen (1999) provides a theoretical explanation of the processes through which children come to recognise illness. She observed the tendency of children to both bring minor ailments—such as a brief
sensation of nausea—to the attention of adults, and also to accept or adapt to more serious ailments, like Cassidee’s disregard of the sore on her finger that morning tea break. These observations are mirrored in the Tūrama School context, where some children present themselves to teachers and the nurses with ‘sore throat’ on a weekly basis, while other children, identified through a class check to have throats so swollen that the nurses wonder how they can breathe or eat, are seemingly unaware of their condition.

While these two things may seem contradictory, Christensen argues that they demonstrate how these children’s understanding of health and illness reflects a process of learning to translate the subjective experience of the ‘incarnate’ body into symptoms and sicknesses according to local cultural classifications and models of body parts and their functioning—the ‘somatic’ body. Christensen draws on the work of Ronald Frankenberg (1980) and Allan Young (1982), who recognise that biological or behavioural signs in the body are interpreted in culturally specific ways and come to be socially recognised as ‘symptoms’ of illness. Christensen adds to this that recognising unusual signs as illness is not a kind of intrinsic ability that people are born with, but that children must learn to identify and distinguish signs of illness from other kinds of uncomfortable or unpleasant experiences of the body. The body is normally very changeable, so actors are not necessarily able to recognise an ‘out of the ordinary’ bodily sensation. As children grow up they come to learn and engage with a complex interpretive and classificatory process of understanding that some bodily sensations of discomfort or pain are accepted as ‘normal’, others are not.

This learning process involves the conversion of bodily sensations (the incarnate body) into symptoms of illness (the somatic body) through several social processes. When children bring feelings, injuries or bodily signs of illness to the attention of adults, the adult will determine whether they are ‘really sick’, often using technical equipment such as thermometers. Once established as sick, adults will administer treatment and separate the child from other children by bringing them to the sick room or keeping them at home. Thus, children link their experience of the incarnate body to its cultural classification firstly through the verification of illness by adult authority, and secondly by the symbolic experience of what happens next: staying home, being cared for, drinking or eating certain things, and having medicine. Notably, for Christensen’s child participants, being ill means ‘then I cannot do, what I usually do’ (1999:187), which could be experienced as annoying, but also signifies their illness as a special event. Finally, Christensen observes children engaging with peers in a ‘collective bricolage of the body’ (1999:210), drawing on their social experiences and
knowledge of the body to link bodily experiences to the ‘somatic’ body. Such a process can be observed in the Tūrama School children’s earlier conversation about the effects of chilli on the body. Thus, peer groups make significant contributions towards children’s understandings of illness, both as sources of information about bodily experiences, and as collaborators in the project of connecting those experiences to cultural models or classification systems.

I suggest that the Tūrama School clinic restructures these processes of linking subjective experiences of discomfort to the diagnosis of illness, resulting in the production of new understandings of illness, and in such a way that children are more likely to present to the clinic to be swabbed. This occurs because the prevention programme is primarily based on a subjective symptom—sore throat—and relies on children identifying when they have a sore throat, either by telling their teacher and presenting at the clinic, or by telling a parent so that they may be taken to the doctor to be swabbed for strep throat. This process presupposes that children have acquired an adult conception of the link between the incarnate experience of a painful throat and the somatic understanding of some forms of this experience as a symptom labelled ‘sore throat’.

In the school, however, children’s somatic understanding of sore throat did not always match the adult biomedical model. Ten-year-old Ngawaina, for example, creates her own terminology to differentiate between her experience of sore throat symptoms and the clinically diagnosed condition that requires treatment:

Ngawaina: I don’t think my throat was… sore sore.

Julie: It wasn’t sore sore.

Ngawaina: I don’t think so.

Julie: What’s the difference between sore sore and not sore sore?

Ngawaina: Sore sore is like… you really need medicine? And not really sore is when… you just need a good night rest?

Compounding the variable understandings of sore throat is a lack of clarity about the difference between sore throat—the symptom—and strep throat as one specific cause of sore throat. Media campaigns focus on sore throat, with little mention of strep throat, while nurses speak of children being ‘positive’, usually without specifying for what. For many children like Ngawaina, sore throat and strep throat become conflated, and sore throat comes to be
understood in somatic terms as its own condition, diagnosed by the clinic staff or the lab. As a result, ten-year-old Mila comes to the clinic not because she feels a painful throat, but ‘so the nurse can check if I have a sore throat’.

In this context children’s ‘illness’ is also being verified by adults in a very explicit and formal way, an institutionalisation of the everyday social process where adults validate children’s illness claims, helping them to link the body incarnate to the somatic body. As in Christensen’s observations of Danish adults and children, the Tūrama School sore throat checks involve adult devices that demonstrate adult competencies and signify illness. Allison will take the children’s temperature, weigh them, look in their mouth with a torch, feel their neck glands and swab their throats. Children pay close attention to the results of these assessments, and in particular the sheet of paper where they are recorded. ‘And every time I come here, she puts me down as a redness and swollen,’ Amberlee (age 10) tells me. ‘Because I watch her, and when she does that it actually kind of freaks me out.’ Importantly, Allison circles ‘sore throat’ on the form to record children’s self-reported symptoms, but for children watching, this confirms their ‘sore throat’ is real. Children therefore do not need the lab results; the assessment, swabbing and recording process is in itself the verification which confirms children’s bodily sensations—whatever they may be—as sore throat. Hence, ‘sore throat’ becomes not only a subjective experience, but a condition which is externally diagnosed.

To further complicate things, sometimes children do not experience a painful throat, but are identified in a class check and swabbed. In such instances, sore throat becomes something that you might not know you have. This experience was very powerful for Mila, who had been ‘given the medicine’ (diagnosed with strep throat) multiple times, but never experienced a sore throat. ‘But sometimes I do have sore throats but I don’t actually feel it?’ she tells me. ‘So, I’ll just think that I’ve got nothing to worry about?’ Because Mila is unable to link her diagnoses with any perceptible symptoms, she presents at the clinic regularly—twelve times this year, she estimated in November—so that Allison can check whether or not she has a sore throat.

Thus the connections children make between subjective sensations of the body incarnate and the illnesses of the somatic body are reconfigured by the school clinic; any symptoms, even an absence of symptoms, can be verified by medical authorities as sore throat. The incarnate experience of sore throat becomes anything ranging from sneezes to lack of feeling, while the somatic category of sore throat shifts from a symptom to a diagnosis. Unlike the interactions
that Christensen observed, children are not simply learning to classify their bodily experiences according to adult concepts of illness, but are creating their own concepts and classifications—sore throat the symptom, and sore sore throat the illness—to explain the inconsistencies between their experience and their diagnosis. This creative coproduction of illness concepts continues in the construction of rheumatic fever.

Children’s knowledge of rheumatic fever

Two shirtless, solemn boys gaze out from a large poster behind Allison’s desk. With a long scar down one of the boys’ chest clearly visible, they oversee the children getting their throats swabbed with the message ‘my brother almost died… it all started with a sore throat’ and the slogan I had come to hear so often: sore throat leads to rheumatic fever, leads to heart damage. Inside or outside of the clinic, I heard children repeat the messages from the poster and from the related television campaigns.

‘Yeah that ad’, said ten-year-old Te Kapua, reciting, ‘my brother almost died. It started with a sore throat.’

Mila tells me how she always likes watching TV, and she likes watching ads, and how ‘that ad always comes up.’ In her narrative though, she adds detail not depicted in the advertisement: a sore throat clinic, a clinic lady, and some medicine which the boy’s mother always gave him:

‘Well, that little boy? Had a sore throat, and told his mum, and the clinic lady, but the clinic lady had nothing to do, so she just gave him medicine. And his mum always gave the medicine to him. And then… suddenly, he started having a sore throat. And like, he couldn’t breathe properly? So he got rushed to hospital, and they had to cut him open. And, and cure what- and cure what happened to his throat. ‘Cause he had heart damage.’

This media campaign (figure 4) was designed by New Zealand’s Health Promotion Agency and includes posters which appear across bus shelters and buildings in targeted areas and a television campaign which has aired since 2014. The campaign was intended to target parents (particularly Māori and Pasifika parents), but also captured the attention of children who identified with the jokey, rugby-playing boys in the video. From what I could tell based on interviews and observation, this campaign formed the primary source of children’s rheumatic fever knowledge; little additional information was formally provided to children, although
Figure 4. Campaign poster for rheumatic fever in the classroom clinic. Poster produced by the Health Promotion Agency of New Zealand. Photograph taken by the author.
Allison would competently answer children’s questions. When the clinic first opened, shortly before I began fieldwork, the clinic staff gave a talk in assembly, but this was not repeated in the time that I was there and few children ever mentioned anything other than the media campaign as a source of their knowledge. An information pack went home to parents with the consent forms, but this was not written at most children’s reading level. The nurse, Deb, gave me a long list of points that she covered with parents when advising of a positive diagnosis and delivering medicine, including storage, adherence, dosage, side effects, reactions, and information for the doctor. When I asked what information Deb gave to the children, the things she listed were reminders—to remind mum and dad to give them the medicine, to bring back their sticker chart—and discussion of their experience of treatment, rather than information.

As a result, children’s detailed and developed conceptualisation of rheumatic fever was constructed using the media campaign as scaffolding. Across the main elements—the sequence of three conditions, the link to death, the primacy of sore throat—children have (to paraphrase both Corsaro (1992) and Geertz (1973)) spun their webs of meaning, using their own experience and what they observed of those around them, and filling in the gaps that adults left blank with knowledges of their own production. Thus, children’s conception of rheumatic fever was consistently unified around a model of its etiology—the most salient aspect. This model was expressed in three stages (or four, with the fourth stage being death), and some children explicitly referred to numbered stages.

During an interview in the clinic classroom after school, three ten and eleven-year-old girls tell me what they know of rheumatic fever. In a collective bricolage reminiscent of Christensen’s description of three Danish children compiling their knowledge of the body’s organs (1999:211–12), they link their experiences and observations of ‘what happens’ and ‘what you do’ to the model of illness presented on the poster. Ruby responds first:

‘Rheumatic fever, ‘cause my cousin had it, and he only made it to the second one, and he almost made it to the heart thing, but they…’ she mumbles, perhaps unclear about what ‘they’ did to prevent heart damage.

‘What do you mean, he only made it to the second one?’ I ask.

Ruby explains, ‘cause like there’s three aye, and you start from sore throat, then you get rheumatic fever, but he never got the things down there.’ She points to the poster behind her.

‘Oh he never got the heart disease’, I say, reading the poster.
As Ngapaea and Alexandra, who have been distracted, tune back into the conversation, I ask again:

‘What do you know about rheumatic fever?’

Ruby begins, ‘It starts from a sore throat-’ She is interrupted by Ngapaea.

‘-Oh it starts from a sore throat, then it goes to heart damage-’

‘-No, it goes to rheumatic fever.’

At this point Alexandra jumps in. ‘It starts with sore throat then rheumatic fever-

‘-and then heart damage-’ all three girls chime.

‘-and then dead.’ Alexandra finishes.

This way of describing rheumatic fever was ubiquitous among these children. Stage one was the sore throat, which, as described above, was represented by a plurality of meanings. Children had a clear idea that if their sore throat was not checked, it could get worse and progress to stage two, rheumatic fever. For most children, stage three, heart damage was linked to death. As the girls explain:

Ngapaea: And the reason we have to have sore throat lady here is, she have to give ‘em medicine to make- you might have heart damage. We get heart damage we’re gonna have to end up in hospital and we have to-

Ruby: -No, ‘cause we can- if- if we too late we can pass away-

Alexandra: -And if we get heart damage we’ll die.

Rheumatic fever was therefore conceptualised primarily in terms of its relationship to sore throat and heart damage, and in its perceived proximity to death, a linear series that would inevitably progress through the stages if no intervention was to occur. As noted earlier, few children included strep throat in their model, reflecting the absence of strep throat in the media campaigns. Consequently, it seems that many children, after receiving antibiotics, may have inferred that they were now at ‘stage two’ of this model and believed they had rheumatic fever. Several children told me about their past history of ‘rheumatic fever’, but described the experience of having their throat swabbed by Allison or a doctor, and taking antibiotics for ten days. ‘Heart damage, I just had that before. For just like two weeks,’ Anton tells me. Some of these children would tell me the doctor told their mother they had rheumatic fever, so it is unclear whether the communication issue was between doctor and
parent, or whether children reinterpreted the information from their parent and attributed their understanding to medical authority.

Yet because children would share their ‘diagnosis’ with their peers, the effect of this was to increase perceptions of the risk of sore throat progressing to rheumatic fever. ‘Jackson’s got rheumatic fever!’ an eleven-year-old girl hissed across the classroom when Jackson was called to the clinic to receive his antibiotics. Children would also commonly attribute the illness or deaths of family members to rheumatic fever. Contrary, however, to how rheumatic fever is represented in the media campaign, child death from acute rheumatic fever is extremely rare, though it is possible that some adult deaths may have represented early mortality from the effects of CRHD. But in many cases, it later transpired the cause of death had been another illness such as pneumonia. Though actual incidences of rheumatic fever remain relatively rare, and most children are probably not even susceptible, as a collective these rheumatic fever narratives created a sense of the disease as much more commonplace, inevitable and deadly than the epidemiological data would suggest. By contrast, the one child I knew of in the senior school who actually did have heart valve damage from a prior bout of rheumatic fever hid his condition from his peers. Nine-year-old Victor would tell his class he was in trouble with the principal when he was called to the office to meet the district health nurse who visited Tūrama School to administer his monthly prophylactic penicillin injections.

While children match their experiences and those of their peers to the model presented in the campaign, the process of connecting the incarnate and somatic bodies was also restructured by the spatial marking of illness. Similar to the children of Christensen’s study, the Tūrama School children associated the experience of being sick with staying home and taking medicine. Though the nurse told me she would try to explain to parents that children only needed to stay home from school if they felt too unwell to go, several children told me they stayed at home for the entire ten day duration of the antibiotics course, despite a lack of symptoms. Ten-year-old Cassidee told me when the ‘doctors’ told her mum she had rheumatic fever they said ‘that I had to stay home for like, two weeks. That was boring.’ Staying at home had the effect of symbolically marking their experience as illness, and children who stayed at home, like Cassidee, were also more likely to be those who interpreted their illness as ‘rheumatic fever’.

In contrast to sore throat, the children’s models of rheumatic fever had less emphasis on symptoms, exactly what the condition is or how it might be experienced. That the most
salient incarnate experience of ‘rheumatic fever’ for Cassidee was boredom makes sense given that any strep throat symptoms probably subsided within a day or two of receiving antibiotics. However, for nine-year-old Marielle, who had given me detailed descriptions of the rheumatic fever campaign, it appears the fear of her diagnosis may have formed her incarnate experience of what she understood to be ‘rheumatic fever’. Even in my interview with her, she gasps between words, unconsciously re-enacting the embodied memory of the illness she associates with breathlessness. ‘I thought I was gonna die because it was my first time,’ she tells me. ‘Rheumatic fever is how… um, you can’t breathe properly? Well you can’t- well you can breathe but you can’t breathe properly, and um, you get heart damage sometimes, if you’re- if you got that very- if you got rheumatic fever too long? And it can be caused by um, sore throats.’

Marielle described the breathlessness as starting after she went to the doctor:

‘…like after. And then I was like, thinking…. what just happened? …. Then I was thinking, like I was gonna be that boy? (in the campaign) Like you know, huh, I was like…’ Her voice shakes, ‘ahhhh, I’ve gotta take my pills!’

The nurse later confirmed to me that strep throat is not associated with breathing difficulties, and Marielle’s breathlessness was most likely anxious hyperventilation, perhaps triggered by the thought that she too would be like the boy on the poster who almost died. Thus while children construct a somatic understanding of rheumatic fever around a model of its etiology, linking campaign information to their social observations and experiences, they also produce a ‘rheumatic fever’ incarnate, primarily experienced as emotions such as boredom and anxiety, and symbolically marked by staying home and taking medicine.

These processes of illness knowledge production therefore shape children’s health practices in several ways. Because of the fear she associates with rheumatic fever, Marielle engages with the clinic more often, and she also becomes vigilant about taking her medicine. Though (with a possible exception being children coping with chronic illness) children’s self-administration of pharmaceuticals is often viewed as inappropriate (Mayall 1993; Christensen 1999), other children besides Marielle also engaged in a variety of semi-independent interpretive practices around pharmaceuticals that contribute to the coproduction of their health. For example, medicine can represent a marker of age status for the children of Tūrama School; because the school nurses administer antibiotics in either liquid or tablet form, children have come to create a shared social understanding that liquid medicine is for young
children, and pills are for older children. This may be why Victor, rather than asking for liquid medicine, spat his pills into the rubbish bin when he struggled to swallow them.

In turn, the self-administration of medicine reflects adult status—something some children may aspire to or be encouraged towards. Nine-year-old Hinemaia, who earns praise from her teacher for her maturity and responsibility, describes helping herself to her brother’s antibiotics from the kitchen when she feels a ‘fever’. The natural extension of enacting that maturity, for Hinemaia, may be taking care of herself pharmaceutically. Likewise, the motherly nine-year-old Teuila, who is Tongan, describes how she diligently takes her antibiotics only every second day so that her younger siblings might have the other half; perhaps an enactment of elder-sisterly care and responsibility.

Children’s conceptions of illness can also inform their pharmaceutical practices. Nine-year-old Dandre and his sister Jordyn are regular visitors to the sore throat clinic, where, they explain in an interview with me, ‘Whaea Allison can check’ to see if they have the ‘sore throat bug’. Across the table littered with empty juice boxes, biscuit packets, paper and pencils, Jordyn draws this ‘bug’ for me, an insect-like creature inhabiting her throat (Figure 5). When I ask Dandre if he ever comes to the clinic when he doesn’t have a sore throat, he tells me ‘Sometimes when… oh, only one time, so they could check if my throat was sore.’

‘What?’

‘See if my throat was sore,’ Dandre repeats.

‘You came here to see if your throat was sore? But you didn’t have a sore throat,’ I clarify.

‘Yeah,’ says Dandre.

After Dandre and Jordyn tell me about how medicine kills the ‘sore bug’, I ask what they think could happen with the bug if they don’t take the medicine.

‘You will die,’ Dandre announces bluntly.

‘Oh!’ I exclaim, surprised.

‘And you get that—’

‘And you get a surgery—’

‘I wouldn’t even like it,’ they clamour over each other.

I wonder if they come to the clinic because they’re worried they might die, and both agree.
‘I thought I was going to die when I didn’t even take it,’ says Jordyn.

‘You thought you were going to die when you didn’t take it…?’

‘Yeah,’ Jordyn explains, ‘because I forgot to take my medicine, and I thought I was about to die. So I took it.’

I clarify; had she just forgotten that day?
‘Yeah, so, so I went to go take it again and again’, Jordyn explained. ‘And then I put my sticker on, and I didn’t even think about it again.’

Dandre and Jordyn’s shared conception of the bug in their throat is the product of nurses referring to bacteria as ‘bugs’ and a media campaign that links the bug with death. However this conception is a powerful driver of their health practices. Dandre, afraid of the bug that will kill you, goes to the clinic so Whaea Allison can check if he has a sore throat. Jordyn, panicking that she might die, takes her antibiotics ‘again and again’, before she puts the sticker on her chart to confirm the medicine has been taken, and she can relax once more.

Both of these children are therefore active participants in the coproduction of their health, but their creative interpretations are also structured by the socio-economic conditions that render them vulnerable to rheumatic fever, a health campaign that supplies them with information about the disease, and a health intervention that medicates children, rather than changing their conditions.

Getting out of class

The high numbers of children attending the Tūrama School clinic can therefore be partially attributed to the restructuring of children’s knowledge production, through which children collectively generate new understandings of illness and the body to guide their practices. They can also be attributed to the fear and anxiety generated by a media campaign that features children and talks about them dying. However, Christensen (1999) also describes how moments of illness forefront the very social notions of privacy, intimacy, weakness and vulnerability associated with feeling unwell. Consequently, illness events accentuate children’s social relations—status, hierarchies, relationships—enacted, for example, in who children notify of an illness event (in Christensen’s case, siblings) and who stays by the sick child while they receive help (close friends). Illness and health care are therefore closely intertwined with children’s peer cultures not only through the production of knowledge, but in the production of relationships.

The situating of the clinic within the school further amplifies the social nature of health care in this context, and this may also help to explain children’s use of the clinic. When I asked about children’s reasons for visiting the clinic, most children said that they went to the clinic only to get their throats checked. Several children also complained that teachers sometimes did not believe their sore throat was real and prevented them from visiting the clinic.
However, as the nurses suspected, some children did suggest that they, or other children went to the clinic to get out of class. This seemed to be connected to Allison’s practice of bringing children to the clinic as a class group, rather than individually, which transformed clinic visits into a social event. Amara, a shy nine-year-old who floated at the edge of peer groups, giggled with me that she ‘sometimes’ wants to get out of class; for her, following the more socially successful children to the clinic may have meant an opportunity to strengthen her relationships. Sometimes, ‘my friends say put your hand up so you can come too,’ Cassidee told me, but on days when she was the only girl who put up her hand, she would visit the clinic by herself at lunchtime for a swab, rather than go with the boys.

The clinic, therefore, could be appropriated by children into the construction of their social relations. However, clinic visits could also offer a moment of respite from the stresses of the classroom environment. A tension between Mrs Randall’s demands for order and quiet as she tried to accomplish the goals of the state, and the subversive resistance of Jackson, Verity, and other children, created a climate of almost constant conflict that appeared to wear on the children as much as it did me. It perhaps is not surprising, therefore, that large groups of children attended the clinic particularly from Mrs Randall’s class (especially since she asked conscientiously every day about sore throats), as well as that of another teacher with a distinctly authoritarian style.

However, talk of getting out of class could perhaps sometimes function as a discourse for children, particularly those with ‘tough’ reputations, to justify going to the clinic to peers. This is where children’s illness knowledge production becomes linked to their social routines; it is not only that children present to the clinic because they are scared they may have a deadly sore throat, or that children only come for the peace and social time. It may also be that children are a little scared of rheumatic fever, but the idea that they visit the clinic to get out of class makes that fear more manageable, and accessing the maternal care of Whaea Allison more socially acceptable.

Glimpses of these complex dynamics arise in my conversation with eleven-year-olds Trystan and Navahn in their classroom after school one day. The pair, who are best friends and ‘cousins’, because their mothers are cousins, have just inhaled my tin of biscuits and Navahn is drawing gang signs with the coloured pens and paper I laid out for them to use.

We began with a brainstorm of the kinds of illnesses they know about, which in their case is extensive.
'My nan had cancer,’ says Trystan. ‘That’s how she died.’

‘My nephew had a asthma attack,’ says Navahn.

‘And I had rheumatic fever,’ says Trystan.

‘And my mum had a epilepsy,’ says Navahn.

They know about pneumonia because Trystan’s auntie just died of it, the class had found out yesterday. She died, Trystan tells me, because she never went to the hospital, because she always drank, and because she didn’t take her meds.

‘Oh. That must have been really hard to watch. Did you know her well?’ I ask.

‘She was like my mum,’ he says. Trystan does not live with his parents.

Rheumatic fever, they know about from the poster in the clinic. ‘It says my brother had-’

‘-rheumatic fever’

‘-ah rheumatic- oh, and it says, it all started from a sore throat,’ Navahn finishes.

‘You guys have like, memorised that poster,’ I comment.

Navahn adds, ‘And one’s got a big as, um- my cousin, his name’s Lazarus, he’s got a big as stitch scar thing there? And that’s what he’s got on the poster.’

The boys estimated they went to clinic about twice a week, the most recent time being that morning. ‘So how come you’re going so often?’ I ask.

‘To get out of class,’ Trystan says, and Navahn agrees, adding that too much running around makes them cough.

But they tell me, in great detail, about how they must finish all their antibiotics, which are called amoxicillin, for 10 days, because there’s 20 pills, so that they won’t get rheumatic fever, because last time Trystan didn’t take all his meds, he was hospitalised, he says for rheumatic fever. And maybe after today Trystan will be taking the medicine again, if his swab from this morning is positive. ‘My nephew went positive 12 times this year,’ Navahn says.

‘Has that scared you a bit?’ I ask them nonchalantly.

‘Yeah.’

‘Yeah.’
‘I might get it again,’ says Trystan.

‘Yeah I’m scared of getting rheumatic fever,’ says Navahn.

‘Are you really?’ I ask, a little surprised at their serious tone.

‘Yeah. I don’t want to go to hospital. I don’t want to die,’ says Navahn. Then he continues, ‘I’d rather kill myself. Nah jokes, jokes!’ He laughs, grabs my recorder and apologises loudly to the speaker, melodrama and mockery steering him away from the edge of vulnerability he has just found himself too far out on.

‘Stink-arse,’ says Trystan.

It is possible that Trystan and Navahn feign a fear of death to cover a well-honed practice of escaping the classroom. But for these boys who live in a world of gangsters and toughness, skipping class may also represent a public excuse to mask a vulnerability which does not fit their staunch social persona. Their expressions in this interview, one moment reflecting on the loss of a parent figure and fear of an illness that seems to saturate their world, the next calling each other names, suggests in themselves an ambivalence about why they are doing what they are doing. If death is an underlying fear turned into a joke, then creating a cultural routine of skipping class also entrenches a socially—and emotionally—acceptable pathway to seeking care.

Conclusion

By December 2016, rheumatic fever rates in Manukau had dropped by a third—a modest success, but not close to the RFPP aim of a two-thirds reduction. The living conditions of families had seen little change, and the problem of child poverty had grown enough to be a primary policy issue in the 2017 elections (Patterson 2017). Meanwhile, Tūrama School children continue to appropriate the clinic processes into their social worlds, while at the same time, the clinic restructures their school day, their knowledge production, and the way they experience their bodies.

This is what is missing from state policy: a view of the embodied, socially embedded practices of children who are collectively carving out a world and their place in it from whatever they encounter. Socialisation through the child’s eyes, as Allison James (2013) writes, sees children as active players in the process of reproducing culture. Child policy interventions, brought to life in context, are transformed through their social enactment, in
interactions between children and the adults who work to care for them. While children’s lives can be structured by adult interventions, children also resist, appropriate, or transform those points of contact with the state through tactical and productive practices that make sense in this umwelt, where classrooms are loud, teachers unpredictable, and death appears close.

This is a view instead of a clinic embedded in social relations and power dynamics: between peers; between adults and children; between those in authority and those who are not. In an institution where authority structures mean children have little power over where they go and who they are with, the clinic may offer a legitimised opportunity for escaping the noise and stress of the classroom. The unusual clinical procedure of examining children in groups also turns health care into a social event, whether children appropriate the clinic as a relationship building opportunity, or establish social routines of ‘getting out of class’ to make accessing care more comfortable or socially acceptable. These social routines then become a new structure guiding children’s engagement with health care.

Generational relations also underpin children’s understandings of health and illness, which are constructed in family and school settings where adults validate children’s sickness, sometimes using technology such as thermometers, and keep children home in bed, or make them go to school (Christensen 1999). When Whaea Allison records children’s self-identified sore throat on the form, she verifies children’s bodily experiences as symptoms of illness, meaning that sneezing can become sore throat, and sore throat can become something you do not know you have. These processes shape children’s patterns of accessing health care in ways that were not predicted by a health policy which does not consider them as significant social actors.

This is Corsaro’s (1992) interpretive reproduction, as described in chapter one: children spinning collective webs of meaning over adult structures and across domains of life. But it is also a generation of culture and practices which are contingent on and constrained by institutional structures and material resources, by the specific social position of children more generally, by childhoods lived in socio-economic contexts more particularly, and by the physical reality of bodies that are small, cold, sick, growing, hot, hungry, tired, disciplined, emotional. Children play an important and powerful role in their health through the way they respond to these circumstances, whether that be constructing polysemic notions of sore throat as simultaneously symptom and diagnosis, taking their antibiotics ‘again and again’ to make up for a missed day, or generating increased perceptions of rheumatic fever risk through their
collective knowledge production and social talk. The processes of coproduction here are therefore the processes that occur between the state and the individual, between structure and agency, between adults and children, and between culture and the body.
In early 2015, just as I was starting fieldwork at Tūrama School, the New Zealand Parliament was debating two members’ bills put forward to amend the 1989 Education Act and expand the social services delivered through schools. The first, known as the ‘Feed the Kids Bill’, provided for government funded breakfasts and lunches in decile 1–2 schools, and was defeated 61–59 at its first reading. Debate on the second, the ‘Food in Schools Bill’ which emphasised teaching life-long nutritional skills, ended in a tied vote, but because a majority is needed, the legislation failed.

The week before the vote on this legislation, I sit on the concrete steps outside a classroom eating my own lunch. As teachers did not eat with their students, I am the only adult amongst all of the senior children, who must sit outside this particular block of classrooms for the first ten minutes of the lunch break, known as ‘lunch-eating time’. This is a somewhat optimistic epithet, as very few of the children are eating anything more than a packet of chips, if that. ‘Where’s your lunch, Caleb?’ I ask the nine-year-old sitting nearby. He shrugs, and gives me
the standard response, ‘I’m not hungry.’ He accepts a handful of cheese-flavoured puffs from Teuila, who sits next to me. This packet of chips is Teuila’s complete lunch and she devours them, tipping the crumbs into her mouth and licking her fingers. A staple food for the children at Tūrama school, packets of chips imported from Asia can be bought from the store around the corner for $1.00 each, making them significantly cheaper than those produced in New Zealand, and, as they come in a bigger packet, an option that can be more liberally shared.

‘Julie, what’s that?’ Teuila asks, looking at my lunch, and I tell her it’s a wrap and the filling is chicken and spinach. I had made it myself that morning. ‘Can I have a piece?’ she asks.

‘A piece?’

‘Of that.’ She points to the spinach.

‘You want a leaf of spinach?’

‘Yeah.’ I let her pick one out. How could I not?

I continued eating, enjoying the sunshine and letting my mind drift away from the conversation. As I finished up my wrap, Teuila interrupted my daydream to ask if she could have the end. Why not, I thought, and handed it over. I didn’t need it, and how do you say no to a hungry child, anyway?

‘Can I have the end tomorrow, too?’

‘Uh…’ I am beginning to see why the teachers don’t eat with the children.

‘Or maybe you could bring me one?’ Teuila tilts her head to the side and gives me a dimpled smile, but I can see in her eyes that she knows what she is asking for is a little shameful.

‘Eh, don’t scab!‘ Teuila!’ Another child admonishes, buying me time to decide how reply. Teuila scowls in response, but with some embarrassment.

‘Hmm, I don’t know. If I brought another lunch for you, then wouldn’t I have to bring everyone lunch? I can’t afford that!’

This prompts an argument about how many lunches I should bring.

‘Yes, bring all of us lunch!’ someone shouts. Other children posit that I should not have to bring any extra lunches.

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1 ‘Scabbing’ refers to the practice of asking or begging for food (or money), seen as shameful or annoying.
‘You could just bring one extra lunch and give it to one child each day,’ suggests Caleb.
‘This morning my dad said tomorrow there would be no lunch. He said we’ve run out of bread. Will you be here tomorrow, Julie?’ he asks hopefully.

This encounter where my lunch was scrutinised, fought over, and shared, was one of many throughout my fieldwork at Tūrama School. While I generally shared when asked, I did manage to solve the dilemma to a certain extent by bringing the most unappealing lunch I could; I discovered that while I liked peanut butter sandwiches on whole-grain bread, the children did not, and my lunch tended to be left alone.2

While I was sharing my lunch with ‘scabbers’, Prime Minister John Key, after an informal phone survey of three schools, argued that the number of children coming to school without lunch is relatively low (Young 2015), and for those who are, Minister of Social Development Paula Bennett invoked ‘parental responsibility’ to defeat the proposed legislation that would have supplied all of the children at Tūrama School with a free lunch (Burrows 2015). The members of parliament who voted against the legislation were strongly criticised by the opposition for being ‘out of touch’ and denying the extent of poverty in Aotearoa.

Meanwhile, the perceptions of children themselves were conspicuously absent from these debates. This invisibilisation of children’s perspectives is routine in child policy, reflecting common assumptions about children as passive recipients of care and provisioning, rather than as social actors who play a role in their own health. Yet, the anthropology and sociology of childhood has repeatedly demonstrated the capacity of children to employ agency in the negotiation of their health (Bluebond-Langner 1978; Clark 2003; Mayall 1993, 1996; Prout 1986), dismantling the notion that adults give care, and children accept it (Hunleth 2017).

Studies of children and care have tended to focus on how children exercise agency in their self-care and care giving (Christensen 1999; Hunleth 2017), but less on the conditions under which children do or do not accept care. Yet, as I will argue, the meanings and practices children construct can mediate the effect of care on their health, so it is important to consider the way that children will interpret state and institutional services.

As apparent in the school lunches debate, children’s health care in Aotearoa is configured around notions of responsibility which shape the particular ways that the state, institutions, parents and children are implicated in the delivery of care and provisioning. However, ideas

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2 This strategy did not, however, relieve the discomfort of doing fieldwork amidst such inequality in my home country. I got to eat my lunch, but I did not feel good about it.
about responsibility for care—who initiates and accounts for what and how care is provided—shift with the positioning of social actors and the state. I have noted the perspectives of teachers of their responsibilities for care of children within the school system in chapter two; in this chapter I contrast the views advocated by the state, often reflected in public discourse, with the views of children themselves. While these perspectives may overlap or, at times, find congruence, they are not the same, and the differences between how the state sees responsibility for children’s care and how this is understood by children themselves can help to explain how and why children engage with different forms of care at school.

Adopting a neoliberal ideology, successive New Zealand governments have discursively allocated responsibility for children’s care to parents, while dispelling any state responsibility for the inequitable conditions underlying poverty and ill health. As a result, state services are designed to ‘support’ and ‘educate’ parents with discharging their responsibilities, while gaps are often patched, with varying degrees of adequacy, by community and corporate organisations. Meanwhile, care in childhood is fundamentally embedded in intergenerational processes, and care relations are a key mode through which social positions are enacted and relationships nurtured. If the state conceptualises care in terms of neoliberal parental responsibility, children see responsibility for care as the enactment of intergenerational and affective relations (Christensen 1999; Hunleth 2017). From the perspective of Tūrama School children, adults, whether in families, schools, or communities, are meant to care for children as the valuable, but vulnerable future of society, while the special care of parents, other whānau, and sometimes even teachers, also forges and strengthens affective bonds. Services which appear universal can hence be accepted as part of the normal social order, just as the state provides free education and primary health care for children. But when a service is delivered in a way that marks out that this care ought to be the responsibility of someone else—a parent—then services that single some children out—such as the charity-provided ‘spare’ lunches at Tūrama School—become stigmatised care. Children’s subsequent engagement with care services at school is therefore the product of an articulation between the state’s view of how responsibility for children’s care should be distributed, and children’s own understandings of how care constitutes their status as children and their relationships with adults.

This intersection speaks to the broader questions of this thesis, as through the particular nature of care services, the neoliberal state provides a structural frame around which children
collectively generate meanings and practices according to the particular ways they understand their social position as children. This relationship is partially captured in Corsaro’s model of interpretive reproduction, whereby children spin spirals of meaning around spokes representing institutional fields (1992, 2015). However, the meanings children make also catch them in their own webs, as they enable and constrain each other’s activities: stigma constrains some food options, while ritual enables others. Meanings which enabled younger children to approach the clinic for care become constraints for older children for whom this care is less acceptable. Children’s cultural production around responsibility and care therefore form new ‘structuring structures’ (Bourdieu 1984), guiding behaviours, patternning bodies, and contributing to the coproduction of their health.

In this chapter I begin with an analysis of how children construct care within generational frameworks, before considering how these constructions mediate two kinds of care service at Tūrama School. Through the example of school lunches and other forms of provisioning, I demonstrate how universalised versus selective forms of care are differently understood within children’s models of adult-child roles and responsibilities. I then briefly return to the rheumatic fever clinic, introduced in chapter four, to explore how such generationally structured interpretations of care shift towards adolescence.

**Care in Childhood**

Issues about care by and for children have been the focus of an extensive body of research in the social sciences. Scholars have largely focussed on challenging Western characterisations of children as passive recipients of care by revealing the range and efficacy of children’s care giving practices, how these are unseen or under-acknowledged by adults, and how adults constrain children’s care practices in different circumstances (Christensen 1999; Hunleth 2013, 2017; Mayall 1993, 1996; Prout 1986; Robson 2004). These debates often centre on practical and ethical questions about the appropriate level of children's caregiving and approaches to supporting children’s care activities, often by emphasising children's agency in order to challenge the Western characterisations of children as helpless, incompetent, and vulnerable which often shape policy and practice. Discussions of street children, for example, disrupt expectations about children's capacities to thrive independently of direct adult care (Davies 2008; Gross, Landfried, and Herman 1996; Mizen and Ofosu-Kusi 2013; Panter-Brick 2002); ethnographies of child domestic and paid labour demonstrate the extent of
children's economic and social contributions (Kramer 2005; Nieuwenhuys 1996; Reynolds 1991); accounts of children's caregiving subvert assumptions about the direction of generational care (Andersen 2012; Hunleth 2017; Robson 2004). Likewise, scholars critique the practices of agencies that intend to help children, particularly in developing countries, often based on Western assumptions of childhood (Hecht 1998; Montgomery 2001). The frameworks, agendas and positions of scholars interested in childhood and care therefore often revolve around tensions between Western discourses of what childhood 'should' be and the realities of life for many children around the world which are not only far from this ideal, but appear to be the ‘choices’ of children themselves.

However the concern with demonstrating children’s agency in care giving has perhaps meant less attention has been paid to how children exercise agency in their care receiving: the conditions under which children will accept or decline care, the meanings children produce that structure how they negotiate care, and how children’s agentive practices mediate the effects of state, institutional, or organisational care on their health. Yet children interact with many forms of biopolitical care which are often subsumed within the public education system, itself a form of state care until transfer of care to legal guardians at the end of the school day. How do children understand, negotiate, and transform those forms of care in the coproduction of their health?

Care, as located in childhood, is grounded in intergenerational processes which shape, as they are shaped by, the help adults and children give to each other. Sociologists interested in the construction of childhood have drawn from Karl Mannheim’s (1952) term ‘generation’ to reference the social structure which constitutes children as a distinct social category, analogous to other structured and structuring categories such as class and gender (Alanen 2001; Mayall 2000; Qvortrup 1994). Leena Alanen (2001) analyses how the ‘generation’ category can be both externally defined, for example by age, or, in the Marxist sense, internally constituted in relation to adulthood (analogous to how class is created through the dependent relationship between wage labour and capital). It is in this latter view of childhood as constructed through intergenerational interactions, where care is implicated as a key mode through which children’s status as children is constituted in relation to adults. How care is understood, and the forms that care may take, of course varies cross-culturally. For example, Jean Hunleth (2017) notes that Zambian children distinguish between Nyanja words for ‘caring’, which translate to protection, help or advocacy, and ‘keeping’, which typically applies to children but also references anyone perceived as dependent, including the elderly
or sick. Thus, an adult may ‘keep’ a child, but not care for them; ‘keeping’ establishes a dependent relationship, child to adult, but ‘care’ creates and maintains interdependent and affective relationships that transcend generations. Children also distinguished between ‘work’, which referred to the domestic or other productive activities they carried out for adults who did not provide material or affective care, and ‘help’, which characterised the exact same activities but only when they reciprocated the ‘help’ children received from adults. Thus for these Zambian children, ‘care’ is defined by the relationships the activities produce, rather than the intrinsic nature of the activities.

Being a Child

For Tūrama School children, however, adult-child relationships are constructed around the direction of responsibility for care; who is meant to care for whom. This came up in interviews, where, when I asked ‘who are the people and what are the places that help to keep you healthy’, children’s answers focussed on adults—particularly mothers, grandparents and aunties—but also fathers, teachers, and Whaea Allison, the ‘sore throat lady’. Children did not name themselves personally, until prompted, and then their contributions were mainly limited to eating fruit. One child named some friends who protected him from bullies. Care, therefore, at least in terms of health, was conceptualised primarily as an adult role.

That adults care for children also became clear when I conducted activities with groups of children asking them to brainstorm ‘what it means to be a child’ and ‘what it means to be an adult’. Children painted a picture of childhood as fun and free, where one can ‘eat lollies’ and ‘don’t need to worry’. This was a particular type of freedom, because although children saw themselves as free from responsibility, they also described a long list of restrictions on what they could do. (Can’t smoke, can’t drink, get told what to do, can’t swear. Kids aren’t allowed to get girlfriends. Kids are not allowed to drive cars. Can’t play violent video games.). Adulthood was described both in terms of responsibilities and of freedom from restriction. ‘Adults have to be more responsible!’ one girl contributed to the brainstorm, and another boy thought ‘(childhood) is better because they don’t get to do the hard stuff. Hard jobs.’ According to the children, adults have to clean, pay the water bill, and ‘be a mother—you have to bath them, change their nappies, feed, buy the clothes, make them a bottle.’ Adults think ‘Man, I wish I never had kids’ and ‘when you grow up as an adult you wish that you were still a kid.’
‘Do you know,’ confided one girl, ‘some people lose their fun, in them? Like they go unfun.’ This dire picture of adulthood was balanced by the perceived freedoms of adults: ‘Adults can go out and go to the movies with all our cousins. And they can go out by themself,’ ‘adults get to go with older people, and they get to play on those-ticket games, the money-pokies!’ ‘Adults can get jobs!’

Most emergent in these brainstorms was the relationship between children and adults; children are cared for by adults. Adults are supposed to help children: ‘when you have a cut they put a plaster on it,’ ‘they make sure you’re fed,’ ‘they love you, care about you,’ ‘they give you lots of ice cream,’ ‘they get to protect you, look after you.’ Adults ‘growl (at children) for not wearing a jumper outside during rain.’

In contrast to many adult views of childhood, children did not consider themselves to be incompetent—just restricted. One group of boys starting phrasing their restrictions in question form, sometimes with answers: ‘Why isn’t children allowed too much sugar? They might get diabetes.’

‘Why isn’t kids allowed to get jobs? Because-kids can’t get jobs because—because they’re at school. They need to learn first.’

‘Why ain’t children allowed to swear? Because it’s naughty words!’

Children also acknowledged their vulnerability, sometimes associating this with their restrictions. Sometimes these vulnerabilities reflected adult fears of violent strangers rather than anything the children had experienced first-hand: ‘Children are little and we can’t go far because we might get robbed or tortured or raped’. All groups of children named vulnerabilities they had experienced as part of being a child, such as being bullied or losing family members. ‘Helpless,’ one boy explained. ‘When you’re getting a hiding. You’re helpless. Especially if it’s from your dad.’

Vulnerabilities also arose from children’s dependency on adults. ‘If you’re young, and your mum and dad die, who would you live with?’ asked one boy who had been living with his grandmother.

In this model of childhood, children’s lack of power and vulnerability is offset by the net benefits of being cared for by parents and being free from the work and responsibilities of adulthood. This is not to suggest that children did not engage in any caregiving activities—of course they did, much like the ‘help’ that Pia Christensen (1999) describes among Danish schoolchildren: taking injured peers to the sickroom, fetching items for parents, giving
affection, making tea. Some at Tūrama School, like Mila in chapter seven, are placed, or place themselves into important caregiving roles for family members. In the Danish context, adults disqualified children’s acts of care, particularly in an institution where the mantra was ‘if you need help, remember to go and find an adult’ (1999:265), while children themselves gave detailed reports of the ways they helped adults. In the case of Tūrama School, it appears children also conceptualise their care giving activities as affective ‘help’; children perform acts of care, but while their autonomy is so restricted, they do not consider themselves to be responsible for care. This may differ in other contexts where children can carry responsibilities with relative independence. From Tūrama School children’s perspectives, the direction of responsibility is what distinguishes children from adults; hence care given by adults to children both constructs and reflects the natural order of things.

Who is responsible for children’s care?

Notions of responsibility also underpin state discourses of children’s care in New Zealand, although the concern here is in establishing just which adults are responsible for children. The neoliberal version of responsibility which couches state views of children’s care is a slippery and multi-faceted concept, in contemporary definitions, indexing ‘individual or collective accountability through judgements of one’s rational capacities, assessments of legal liabilities and notions of moral blame’ (Trnka and Trundle 2017:4) and connoting agency and autonomy, but also discipline and obligation (Shore 2017). Though the notion of the responsible citizen is not new, in its contemporary form personal responsibility represents a key mechanism of neoliberal governmentality (Rose 2007). When applied to health, responsibility discourses enact a form of biopower (Foucault 1990) which mobilises independent citizens to govern their own bodies using a rhetoric of ‘personal choice’ supported by ‘health promotion’ education rather than government or doctor authority.

This contemporary personal responsibility, as invoked by Paula Bennett (Minister for Social Development) in the school lunch debate and understood by parents and teachers, reflects a shift in the last few decades from New Zealand’s historical relationships between individuals, society and the state. As detailed in chapter two, social policy in Aotearoa was traditionally based on the accepted principles of the 1930s welfare state, with a wide consensus for a collective responsibility ‘in order to protect the unfortunate and safeguard the nation’s children’ (Rice 1996:483). In the past, the State also took an active role in supporting
children’s nutrition, including government provision of milk in schools between 1937 and 1967 (New Zealand Milk Board 1978). However, the changing family structures of the 1980s—declining marriage, increasing divorce rates, and a significant rise in single-parent families—led to increased demand on welfare services and spending. The subsequent decline in support for the welfare state among New Zealand people coincided with similar trends in other Western countries and the growing dominance of neoliberal policies: economic deregulation, a reduction and commercialisation of state services, and the repositioning of responsibility for wellbeing with individuals, their families and local community (Nairn, Higgins, and Sligo 2012). Within this ideology, children are generally subsumed under the umbrella of familial responsibility (Trnka 2017), and are invoked in powerful discourses that morally frame the good parent as one who ‘takes responsibility’, as seen in the school lunches debate, as well as one who reproduces responsibility by raising responsible citizens. Yet the neoliberal version of responsibility sits uncomfortably as a framework for lives that are constrained by inequities in power and limited choice, and for children, who cannot be held responsible for themselves. Problems arise when parents do not, or cannot meet children’s needs; when children are coming to school without lunch, or suffering from health problems stemming from poverty. A deficit-framed popular discourse of personal responsibility blames parents for causing these issues, meanwhile, what responsibility does society have to the children?

The answer, for New Zealand, has been an uncoordinated patchwork of services to plug the gap. For Tūrama School children without lunch, three services offer alternatives to going hungry. Firstly, neoliberal reform has seen a stepping back of state welfare services in favour of ‘education’ to equip citizens to take responsibility for themselves. Under the mantra of ‘health promotion’, therefore, the 2005 Fruit in Schools programme (described in chapter two) offers free fruit three times a week to children in low decile schools with a mandate to promote healthy eating (Boyd 2011). Secondly, the ideology of the welfare state has been replaced with a language of ‘community empowerment’, encouraging local organisations and NGOs to step forward as the state withdraws and placing the decision-making—and the responsibility—for health onto local communities (Cushman 2008). Thus when Labour MP David Shearer, who put forward the “Feed the kids” bill, stated that he had changed his thinking and that “each school community should be resourced to find and deliver its own long-term food solutions” (Young 2015), he was likely influenced by this popularised notion of deflecting responsibility back onto communities, leaving the decision-making and
implementation of necessary programmes to schools, NGOs, and community groups. As part of this community effort, children who come to Tūrama School without lunch have the option of a ‘spare lunch’ provided by charity organisation KidsCan (www.kidscan.org.nz). Finally, as part of a global business trend, the socially responsible corporation may claim ‘responsibility’ for helping to combat poverty and thus partially releases the state from pressure to intervene. In the case of school lunches, this has taken several forms, including the marketing strategy of catering company Eat My Lunch which promises to donate a free lunch to a child in need with every lunch purchased (Tūrama School did not take part in this programme at the time of my fieldwork), and on a grander scale, the introduction of the Milk for Schools programme by dairy co-operative Fonterra (www.fonterramilkforschools.com).

In theory, these three levels of intervention should mean that every child eats a substantial meal at lunchtime. Yet the efficacy of these services depends on how they are viewed, experienced and interpreted by children, for whom care takes special meaning in the constitution of identities and relationships. Rather than passive recipients of whatever charity is offered, children actively make meaning around these provisions which influence when and how they are accepted. Children’s nutritional status is therefore a coproduction between the neoliberal governance that shapes their food options, and the social meanings and practices that children collectively generate around them.

Pōhara, or ‘not hungry’

There is a distinct dietary pattern across children at Tūrama School. Until they get home from school, standard fare for a child at Tūrama School consists of no breakfast, chips or biscuits at lunchtime, a carton of milk and a piece of fruit. Although this is a typical lunch, there is some variation. Some children, if they have been given money, will buy some hot chips or a pie for breakfast on the way to school. A minority will request a ‘spare lunch’ and receive a sandwich and snack supplied by the KidsCan charity. Parents are more likely to supply younger children with a packed lunch, but often expect the children I worked with, aged 8–12, to arrange their own lunch.

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3 As meagre as this sounds, the school nurses, weighing children with sore throats to determine their antibiotic dosage, noticed that children’s weights tended to be lower after the school holidays. The nurses suggested that the eating routines built into the school day ensured children were typically eating more at school than they would at home.
This variation may contribute to or reflect some of the different processes through which malnutrition is produced in this context; children’s body sizes tend to bifurcate into the very underweight and the very overweight, both manifestations of malnutrition (Ministry of Health 2012; Oliver et al. 2011; Utter et al. 2007).

This signature diet comes partly as the result of both deprivation and intervention; as parents struggle to supply lunch for their children, alternatives are offered through school services. However, children translate these services through a lens of adult-child roles and responsibilities, and the resulting social meanings influence which provisions children will tend to accept. Children’s nutritional status is therefore not only the product of deprivation and intervention, but their own collective social practices. Children’s understandings of poverty, assistance and adult-child roles mean that while the milk and fruit are typically accepted, most children don’t take up the offer of the spare lunch, even when they have little to no food of their own. Instead, most children without lunch will claim they are ‘not hungry’, rather than ask for a spare lunch.

The ‘spare lunches’ provided at Tūrama School consist of a sandwich—usually peanut butter, occasionally jam—on white bread, which is often supplemented with a muesli bar or small packet of nuts as donated stocks allow. The sandwiches are made in bulk, using donated ingredients, by a group of volunteers in the school kitchen every Tuesday, wrapped in cling film and frozen. Each morning, teachers are supposed to ask their class who needs a spare lunch, and record the names of those who put up their hands. The numbers are sent to the school office, and the secretary will remove the required number of sandwiches from the freezer, with the expectation that by lunchtime they will have defrosted. The spare lunches are not popular; one child described them to me as ‘cold bread’, and many children claim a distaste for peanut butter (which I capitalised on to avoid sharing my lunch). Staff estimated that they gave out around 20 spare lunches a day, but noted that it varied; the day before payday more were needed, and some individual teachers would push spare lunches more than others at various times, though this may not have much effect on how many spare lunches were actually consumed.

One morning, while sitting on the mat with a class who I had not visited recently, I am surprised when after calling the roll the teacher, Mrs Charles, instructs the class to go and get their lunches. Some children get up and go immediately to the corridor where their bags are

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4 Their dislike for peanut butter could well be because of its almost invariable presence in spare lunches.
hung, but others hold back, avoiding eye contact. As they return to the circle with their lunches in front of them I get the sense this has been happening every morning for at least a few days.

As the stamping feet and chatter dies down, Mrs Charles’ voice, quiet and low, cuts across the classroom. ‘Right. Can you tell Julie why we are doing this?’

‘Because you need to have lunch or your brain will fall asleep,’ one of the girls replies.

‘That’s right,’ Mrs Charles confirms in a sing-song tone. She then proceeds to inspect each child’s lunch, one by one. Mila has only a muesli bar, so Mrs Charles writes her name on the spare lunch order form. ‘Where’s your sandwich, Ruby?’ she asks. ‘You usually have a sandwich. You make it yourself.’ Ruby, sitting with a packet of chips and an orange (the only piece of fruit in the class), mumbles and avoids eye contact. Many children have only chips or biscuits, and so their names are recorded. Five or six children, mostly boys, do not have anything in front of them. Some of them say ‘my mum is bringing it later’ and Mrs Charles responds, ‘I know what that means!’ and writes their name down too.

When the lunch bell rings, a bread bag stuffed with the ordered sandwiches arrives. Mrs Charles looks at the order form and calls out names. Mila! Ruby! Kauri! Wiremu! Come and get your lunch! One or two children come to claim a lunch, but most of the class have already disappeared out of the classroom, leaving Mrs Charles holding the bag of half-thawed sandwiches.

I watched Mrs Charles repeat this process a few days later, but she did not continue with it much longer after that, later telling me:

‘It got to the stage where I was being the food Nazi, and it got to the stage where I was putting them on the spot, and they would order the spare lunch and then they would just chuck the spare lunch, they would just chuck it away.’

While children, when I asked why they did not order a spare lunch, claimed they weren’t hungry, in interviews after school I was able to gain more insight.

Twelve-year-old Arya told me how in her class, ‘Hardly anyone gets spare lunch ‘cause they’re shamed? ‘Cause it’s like, they’re pōharas? [poor]’ She continues, ‘Today- someone pranked my name on the spare lunch. And I know, that’s not my handwriting.’

‘Have you ever wanted to get a spare lunch but you haven’t because you didn’t want people to make fun of you?’ I wonder.
‘No,’ says eleven-year-old Nikora, who is with us.

‘We always have lunch, aye?’ Says Arya.

‘You always have lunch,’ I repeat.

‘We never, ever, ever- oh well, in term one, ‘cause my mum brought like things that, should not be… she bought… phones, and stuff.’

Having established that her lack of lunch had been due to what she framed as her mother’s irresponsible purchasing, Arya goes on to tell me how much her Dad gets on payday, which to her is a lot. ‘I can buy heaps of food if got that much. But they get smokes and that’ she laughs, embarrassed, again referring to how her parents spend their money, rather than not having enough. ‘They could like donate to… Save the Child and stuff.’

I am not sure what she means, but Arya explains that her mother had donated fifty dollars to buy food for African children.

‘But were you saying that in term one you weren’t getting the right kind of lunch?’

‘Yeah. My- I had… to go to school with no lunch.’ Arya laughs, to show this didn’t upset her. ‘But I never got spare lunch, I just like ate the fruit and that. But in term two, and now, I get money and stuff.’

In Patricia Grace’s (1986) novel Potiki, the Māori people reclaiming their land speak of being, of course pōhara, but also emphatically not pōhara, this time referring to the poverty that comes with the cultural death of a people. I understand the word pōhara to connote the indignity and disenfranchisement of poverty, more than just the material state of deprivation, and I suspect it is with this valence that Tūrama School children use the word to signify the absence of something greater than lunch. In this conversation, Arya casts those who get spare lunches as pōhara, but positions herself outside of this category because she always has lunch. Then remembering an earlier time when she didn’t have lunch, she blames her mother for irresponsible purchasing, rather than accept the label of pōhara herself. It is not that her mother did not care, because she never took a spare lunch, like those other, pōhara children. Arya then details her parents’ finances to show they were definitely not pōhara, and tells me about how her parents donate to Save the Children, contrasting her family with the children in Africa who really don’t have food. Now, she tells me, she always has lunch, because her mother gives her money.

‘So do you buy some lunch on the way to school?’
‘Yeah. Today I bought some lollies.’

To Arya the lollies she buys count as lunch, so she doesn’t need to consider a spare lunch, and she clearly isn’t pōhara. She continues:

‘That’s my lunch and… and I bought a packet of chips.’ She tells me her brother stole her chips to give to his friends, and her voice drops, almost to a whisper. ‘That was for my lunch. That’s why I have nothing besides milk and fruit. Oh no, just milk.’

Because today was not a fruit day, her lunch has consisted of milk and lollies. Yet this is still a socially acceptable meal. For these children, lunch means more than sustenance; it is also a highly visible symbol of socio-economic status, and of a parent who is giving care as parents should. Because the spare lunches require special—and public—request, this well-intended charity service marks out children who have unfulfilled needs, establishing the idea that lunches are not meant to be supplied by the school. Lunches are meant to be supplied by parents. Mrs Charles’ regime of publically identifying whose parents have and have not supplied them with sufficient lunch, while unusual and short-lived, represents an extreme reinforcement of these social meanings. The unappealing nature of ‘cold bread’ further reinforces this as substandard, and therefore stigmatised care. Thus neoliberal discourses of parental responsibility are translated into the nature of service provision, which children in turn reproduce. Echoing public discourse, children, too, can blame their parents for a failure of responsibility rather than a lack of income.

This invoking of parental irresponsibility appeared in conversations with other children also. In an interview with a group of girls from Mrs Charles’ class, Ngapaea says to Ruby, ‘You mock Tāmati when he has to get spare lunch.’ As the girls dissolve into giggles, Ngapaea explains, ‘she mocks Tāmati when he gets spare lunch. Because he like, got nearly- he got like spare lunch for like a whole term.’ The girls shriek with giggles.

‘Why do you think some children are getting spare lunch?’ I ask.

‘Because we got a day with no lunch,’ says Ruby, meaning that she, and others sometimes don’t have lunch on a given day.

Ngapaea is not willing to claim membership of this group. ‘Or, their parents um, are broke.’ ‘Usually they spend it on beers, and cigarettes’ adds Alexandra.

Like Arya, the girls here emphasise that getting a spare lunch does not imply a family who has limited economic means, but specifically a parent who has misspent their money, prioritising beers and cigarettes. Also like Arya, they position themselves as different from
those children. Ruby claims she’s never had spare lunch, and Ngapaea admits she had spare lunch when she was younger, but not anymore ‘because my mum is organised with the lunch’.

‘Yeah that’s like us,’ says Ruby. She has forgotten—or is not willing to remember—the two occasions I observed Mrs Charles ordering her a sandwich after lunch inspection.

‘Ok, so you always have lunch now. And so now you can just laugh at Tāmati for…’ the girls giggle. ‘You guys are so mean!’

‘He laughs at us!’ Ngapaea protests.

‘Cause we be naughty, like we talk…’

‘We get in trouble ‘cause we talk in class, and then- shame!’

‘But like, when he orders spare lunch we like…’ The girls laugh.

Ngapaea finishes, ‘But he just chucks it away.’

In the social meaning collectively understood by children at Tūrama School, asking for a spare lunch designates a child as pōhara. For times when they do not have lunch, then, children have created a range of strategies to negotiate this identity. The most common and socially accepted strategy is to claim ‘not hungry’, making choice from necessity (Bourdieu 1984), a state that is not only discursive but may become embodied, as I describe in chapter six. Some children take a spare lunch and either accept the lower social status that comes with it, or, like Tāmati, throw the lunch away under pressure from teasing. Some children, like Arya and Ngapaea, appropriate the language of responsibility to reframe their lack of lunch as disorganised parenting so as to avoid the pōhara identity. Finally, at times the school receives donations of a food item popular with children, such as fruit cups or a packet of trail mix. When this high-value product is included in the spare lunch, some groups of children will collectively decide it is acceptable to order a spare lunch in order to obtain this item. In these circumstances, while the group ostensibly orders a spare lunch solely to eat the chocolate out of the trail mix, it is possible for individuals to make a performance out of picking out the chocolate, and then quietly consume the rest as well.

Possibly the most significant effect of these dynamics, however, is to transform definitions of lunch. When having no lunch designates a child as pōhara, any amount of food, no matter how insubstantial, is enough to be classified as ‘having lunch’, symbolically marking both economic means and a parent’s care. The pattern of nutrition at Tūrama School therefore is
founded in the one dollar packets of imported chips and fifty-cent lolly mixes available at the corner store on the way to school.

Milk

The social meanings children ascribe to spare lunches, constituted through notions of who should be supplying the lunch, therefore limit the acceptance and consumption of those lunches. Yet the supplying of food by adults other than parents is not in and of itself interpreted as stigmatised care. Other forms of provisions through school services are taken up with enthusiasm. Children from each class are given the task of collecting the fruit from the Fruit in Schools programme three times a week and this is distributed indiscriminately to everyone at the beginning of break time or eaten together during class time. Likewise, Fonterra’s milk has now been knitted into the daily routine of Tūrama School.

The ‘Milk in Schools’ programme was initiated in 2012, amongst the growing debate about child poverty and the role of government and schools in children’s nutrition. Applauded by the government and media alike, Fonterra distributes a free 200ml carton of reduced fat milk to 145,000 children at any primary school which chooses to participate: currently 70% of New Zealand’s schools (‘Fonterra Milk for Schools’). As well as invoking a nostalgia for the historical (state funded) provision of milk to schoolchildren between 1937-1967 (New Zealand Milk Board 1978), the near-universal embracing of the programme is probably at least partially because Fonterra, New Zealand’s largest company and a national icon of economic success, is owned by 13,000 farmers, and is thus emblematic of the idealised, hard-working, tax-paying, rural Kiwi rather than corporate self-interest. Fonterra’s justification as outlined on their website—to give every child “access to dairy nutrition every school day”—does not mention the opportunity to market milk to children, entrenching their produce as an institutionalised part of childhoods, or the clear commercial benefit of a generation of children associating milk with nutrition and accustomed to having milk as a daily part of their diet. As milk is marketed as a highly-nutritious food, the promotion of their product to children is seen as a socially responsible thing to do—a sentiment reflected in sponsored media coverage (New Zealand Herald 2016).

In contrast to spare lunches, at Tūrama School, nearly all children drink this milk, and may even steal or squirrel away additional cartons of milk for later. Thus milk has in recent times
become the staple food of Tūrama School children, largely because of the way it has become embedded in a collective social ritual of consumption.

On a typical day in Mrs Randall’s class, we come into the classroom after morning tea, breathless and sweaty from our soccer game. I claim a spot on the mat with my back against a cupboard, and Cassidee slips in beside me on one side, Caleb on the other. The rest of the children arrange themselves into a circle. Usually when we sit on the mat in Mrs Randall’s class, we sit facing the teacher at the front for instruction. But every day after morning tea the class come in and sit in a circle on the mat. This is symbolic of milk time, the one time that we can see everyone in the class across the circle.

Mrs Randall opens the box of milk and hands cartons to the child next to her, who passes them on until the milk has made a full circuit and all of us have one. The cartons are cold, as Mrs Randall has installed a small fridge in the classroom to keep them in, and some of the boys across the circle hold the cartons against their red cheeks to cool down after soccer.

We shake a nd puncture our cartons with our straws and sip slowly, prolonging this time of relaxation, because it is a peaceful social time. Today is Whetu’s birthday, so Mrs Randall hands around a container of chocolate biscuits to celebrate. ‘You can have one of each kind,’ she instructs. Some children have brought biscuits from home as well, so they can have ‘milk ‘n cookies’.

‘Miss, can we have our banana now too?’ asks Dandre. ‘So we can have banana milkshakes!’ Mrs Randall laughs, reaching for the basket, and the bananas—the fruit of the day for the Fruit in Schools programme—get passed around as well.

Mrs Randall is drinking a milk too, and she leans back in her chair and chats to Amberlee about her new baby sister. We relax and talk amongst ourselves, but with mouths full of straws and banana and biscuit the chatter is not overpowering or loud. Finishing her conversation, Mrs Randall opens her bookmarked copy of ‘Matilda’ and starts reading aloud.

I sip my milk slowly as I listen. It does not taste bad, but is not something I would normally choose to drink. At the start of the year I used to decline the offer of milk, and sometimes bring an alternative, but I soon found myself drinking the milk for the pleasure of being included in the group. The milk drinking ritual brought the class together in companionship and bonding as they are literally and symbolically nurtured by their teacher and each other. The children do not have to drink the milk, but they usually all do. One time Dandre, upset after being told off, refused to take his milk and his banana. He sat in stony silence while the
children around him alternated between drinking their own milk and asking if he was sure he didn’t want any. Usually delighted for ‘banana milkshake’ day, Dandre’s rejection was of the social bonding more than of the food, an expression of his upset state.

After we drain our cartons, making sucking noises to get the drops in the corners, we fold them according to the method prescribed by the recycling guidelines. The children taught me how to do this at the beginning of the year, and it took me a couple of tries to learn, but now they say I am very good at it. My carton is always very neat and flat. It requires a bit of technique but the children are very adept at it, and often a child will ask if they can do mine. Much like folding origami or paper aeroplanes, the folding of the cartons is an activity where we can demonstrate skill, and now we fold juice and other cartons in the same way. One of the boys spills the milk all over himself as he folds the carton. ‘That’s why you’re supposed to drink it dry,’ says Mrs Randall. A few of the children begin to chant ‘drink it dry, fold it flat, send it back!’ Others join in the chorus as we take our folded cartons outside to a special-purpose bin, which will later be emptied and collected by Fonterra for recycling. The cartons are shipped to Thailand to be made into roofing tiles.

This routine is similarly embedded into life in other classrooms. But one day in Mrs Charles’ class we were sitting in our circle after morning tea, and the children ask, ‘where’s the milk?’ ‘Where is the milk!’ says Mrs Charles, and she asks one of the children to run and find out where it is. We sit, looking at each other across the circle, waiting. There is a knock at the door and another teacher puts her head in. ‘Do you know where the milk is?’ she asks. ‘All my kids are ready for their milk!’ As Mrs Charles gets up to discuss where the glitch in the delivery system may be, I laugh ‘this school falls apart when the milk is not delivered, huh?’ ‘Yeah’, says Ruby. ‘After morning tea we feel like milk, and there’s no milk.’

This deep routinisation of milk drinking—highlighted on the day the milk did not arrive—contrasts with the stigma endured by children who requested a spare lunch. Instead, children’s milk consumption is not just because it is provided, but because its universal distribution democratises its consumption, and because the ritual practices collectively generated by teachers and children together—the circle, the cookies and banana milkshakes, the carton folding—make milk more about the coproduction of shared social experience of togetherness than nutrition. Spare lunches, which require children to single themselves out, and thus indicate difference, poverty, or parental irresponsibility, are less accepted, when
chips and lollies, bought cheaply, can symbolically represent material assets and a parent’s care.

The pattern of nutrition at lunchtime at Tūrama School therefore is not only a product of deprivation and intervention but also of children’s own cultural meanings and social practices which are collectively produced and directly shape children’s bodies in patterned ways. A diet of lollies, bananas and milk has some nutritional value, but is far from a balanced diet and high in sugar. Furthermore, though a normative discourse of milk as a growth food for children is used as a global marketing strategy (and rarely questioned), Wiley (2007, 2014) argues that the correlations between milk and growth do not represent anything intrinsic to milk, but simply reflect the protein and caloric content of milk—though milk can be a convenient method of delivery. Furthermore, milk comes with its own set of digestive problems. While onset of progressive symptoms for individuals with lactase non-persistence genes typically occurs around adolescence or early adulthood, it is common for children to develop symptoms, in some populations before the age of five (Heyman 2006). Lactose intolerance is common among people of Polynesian ancestry; in a sample of 160 adults with self-reported Māori or Pacific ancestry, 34% had the genotype for lactase non-persistence, and the frequency rose with increasing Polynesian ancestry (Roberts, Merriman, and Upton 2010). Yet when the government abdicates responsibility for supplying nutrition, they also leave it in the hands of corporations to decide what form that nutrition will take—a decision that will not be based on objective evaluation of options.

The specific forms that Tūrama School children’s food options take therefore are a product of much wider economic and political circumstances, and particularly ideological notions of how responsibility for children’s care should be distributed. Yet children also actively negotiate, accept or decline these options, based on different moral interpretations of different forms of provisioning. Tūrama School children are not aware that only schools in the poorest communities receive fruit, or that for some schools in wealthier areas, the free milk is deemed not worth the programme’s administrative burden. Because both of these forms of provisioning are universal and routinisied, their supply is part of the normal order of things; it is the role of adults in general, including schools and teachers, to care for children. The confluence of neoliberal ideologies of responsibility for children’s care, and children’s own perceptions of the roles of adults, therefore creates the conditions for practices which affect children’s health in distinct ways.
Sore Throat Clinic

Thus the neoliberal version of responsibility comes to shape children’s health practices via the particular forms that care services take, contributing to the avoidance of spare lunches in contrast to the uptake of milk. Ideas about the distribution of responsibility also inform the Rheumatic Fever Prevention Programme, which in turn creates particular experiences that children, again, interpret within intergenerational structures.

While neoliberalism has restructured health policy according to notions of personal responsibility, the movement has also transformed the construction and governance of childhood (Tap 2007), making it what Rose calls ‘the most intensely governed sector of personal existence’ (Rose 1999:123). The surveillance medicine that emerged in the early 20th century targeted the child for intensive monitoring of their growth and development through clinics and schools (Armstrong 1983), but neoliberalism has reframed this surveillance in terms of personal—and in the case of children, parental—responsibility for monitoring health.

The way the state views responsibility for rheumatic fever, therefore, means that Tūrama School children still live in poverty, but get a clinic, preventative antibiotic treatment, and a campaign designed to remind their parents to take action for sore throats. An archetypal example of surveillance medicine, the intervention programme involves both screening and health promotion and thus brings illness out of the hospital and into community life, constituted as a constant risk to be monitored, but conceptualised, prevented and treated at the level of the individual family. Responsibility for preventing rheumatic fever is placed on the shoulders of families, starting with media campaigns urging parents to take their child to the doctor for a sore throat. ‘If we got it checked earlier, he wouldn’t have this,’ a young boy says in the television advertisement, pointing to his brother’s scar. ‘If I knew that this sickness started with a sore throat, I would have taken him to the doctor earlier,’ says his mother in another advert. ‘It’s not cool to take the risk of “I’ll see if it’s okay tomorrow.” As soon as your kid gets a sore throat, you need to get it checked,’ warns the mother of another young boy. These statements, coupled with horrified descriptions of having a child in hospital and seeing them almost die, give the implicit message that the family is responsible for

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5 The programme is not officially classified as medical ‘screening’—a label that brings with it particular ethical standards due to the increased anxiety screening produces—because only children who are identified as showing symptoms are checked. I would argue that class checks and teacher questioning to identify symptoms constitute screening, if not by medical definitions, then at least in effect.
rheumatic fever prevention. There is no mention of the fact that between one and two thirds of acute rheumatic fever cases are not preceded with a known sore throat event (Robin et al. 2013; Veasy et al. 1987), or that children themselves may not recognise or report their sore throats. The framing of rheumatic fever prevention does not recognise how impractical it may be for families with multiple children and working parents to take children to a doctor for every sore throat—though this issue is at least partially solved for communities with school clinics—or that outbreaks of rheumatic fever might point to hazardous socio-economic inequality and be better addressed through economic policy. The illness is characterised as an individual problem, the solution is for families to monitor and medicate their children, and the school clinic is established to support parents and children to deal with their problem. Thus, out of ideas about responsibility, rheumatic fever enters the lives of all children at Tūrama School to stamp their bodies with antibiotics, anxiety and hypervigilance of the throat; an embodiment of surveillance that I explore in chapter six. Here, I consider the way this state conceptualisation sits alongside children’s views on responsibility for care, shaping the way that children accept with enthusiasm, avoid, or negotiate care from the clinic.

In many ways, the integration of the sore throat clinic into Tūrama School has had positive effects. As I described in chapter four, large numbers of children access care from the clinic on a regular basis, and have a good experience of health care that many translate to a positive view of external health care services as well. Though primarily a sore throat and skin infection clinic, Whaea Allison and Deb do identify and treat or refer children for other illnesses on an ad hoc basis. Children’s enthusiasm, however, drops off as they graduate to the combined year 7/8 class, aged 11–13. Whaea Allison had to make special effort to make sure these older children—particularly the year eights—were checked regularly. The clinic therefore works very well for children, but less well for adolescents.

From the perspectives of children, the clinic is not stigmatised care; like the milk, the clinic intervention is universal—every child is checked under the throat swabbing programme, not just those whose parents cannot get them to a doctor. The clinic system also fits with children’s own model of childhood, reinforcing the adult-child relationship of care. For children, adult monitoring of their health status is a normal care practice that establishes adult and child identities. Surveillance medicine, therefore, is another form of adult governance over children’s lives (Rose 1999) that this generation have grown up experiencing as the normal order of things. Teachers ask children if they have a sore throat in the same way that mothers check on children’s health. Whaea Allison is warm, motherly and affirming, and
takes children through a diagnostic process that is translated into a practice of care. Ten-year-old Tupono described this process in great detail for me:

‘First of all you stand on the scales. They check your weight. And, sometimes that could do with sore- like a illness you can put on heaps of weight or you could lose heaps of weight? And… so I stayed the same, oh- thirty to thirty four? A lot less than everybody in my class and… then next they get you to check your temperature and your ear with the ear temperature thingy. What is it called? It’s a m… Do you know what it’s called?’

‘Thermometer?’ I supply.

‘Oh yeah’, Tupono continues. ‘I knew it had ‘meter’ at the end. Then they get this, so you know those things that you get your ear wax out, those pokey things, they have a long one but not double si… Do you know what it’s called?’

Through this process, children understand Whaea Allison to be assessing their sickness, and in doing so she is showing care and protecting them from the dangers of disease. Tupono, a bright boy with a particular interest in medicine that he has acquired through reading medical literature in church, gave the most technical description of the clinic processes of all the children I talked to, including an interpretation of why he was being weighed. Children were actually weighed to determine the dosage of antibiotics, but often took careful notes of their measurement and connected this with the monitoring of their health and development: ten-year-old Ngawaina imagines Whaea Allison weighs her ‘to see how big I’ve grown.’

Meanwhile, some children mistook Allison’s assessments for treatment. I observed a young girl, after being swabbed, saying she still felt sick, and Allison gently explained, twice, checking for understanding, that the swab is a test, not a treatment. When she was not made aware that children thought they were seeking treatment, Allison was not able to clarify. Ten-year-old Anton, a regular visitor to the clinic, described how he would feel better and his cough would be gone for the rest of the day after being checked at the clinic; he understood ‘sore throat’ to mean ‘coughing and fevers’ and connected the abatement of his cough with being swabbed. He did note that the cough would be back the next day though, but he was only allowed to go to the clinic one day every week:

‘…if I get checked, like my coughs is gone, for like the whole day? And then after the other day, I can’t go to the sore throat lady, because you only allowed one day, and then… it just comes back.’

Hence, children’s experience of the clinic processes matches and reinforces their concept of adulthood and childhood, that children are cared for and protected by adults. Children may
come to the clinic seeking the care and approval of Whaea Allison as much as to have a sore
throat checked.

For older children, however, these dynamics can have the opposite effect. Children in the
combined year 7/8 class, aged 11–13, presented less often to the clinic than those in years 5–
6; they drag their feet to avoid class checks, and Whaea Allison must make a special point of
tracking these children down. This form of surveillance medicine, configured around adult-
child dichotomies, is welcomed by those who embrace their child status, but resisted by older
children who see themselves transitioning towards adulthood and anticipate adult relations
with health professionals in negotiating their care. The clinic procedures, however, position
them squarely as children. At a time when young adolescents are claiming their bodily
autonomy, the clinic requires them to hand their bodies over to an adult to be weighed,
inspected, felt, and inserted with thermometers and swabs. The public nature of the clinic’s
work, which younger children capitalised on as an opportunity for social interaction, became
an embarrassing intrusion into privacy for older children. Sometimes older children would
instead visit the clinic quietly at lunchtime, when their check could be made discreetly and in
private. Furthermore, even the year six girls had begun to become self-conscious about being
weighed: ‘Today my weight was 51,’ Ngawaina whispered to me in our interview. ‘I gained
heaps of weight.’ By year seven or eight, for many girls, in particular, weight is no longer
positively associated with growth, and the process of being weighed put them into a
vulnerable and embarrassing position.

Children’s engagement with care services is therefore mediated by their conceptions of
adulthood and childhood and how their perceive their own status within this binary. In a
model where childhood is established in relation to adulthood, the shifting and liminal stage
of adolescence makes for unstable identities within a framework of adult authority and child
vulnerability, adult care and child help. These young people often still want or need care;
they are still disturbed or frightened by the rheumatic fever campaign, and some do negotiate
their own path to the clinic’s care. Despite its common association with childhood, they still
drink their milk together, long limbs sprawled on the mat, while their teacher reads them
Māori myths from a picture book. But the children who in year six were still taking a spare
lunch, never do as year sevens and eights, and their teacher does not bother asking any more.
These are young people at the beginning of what will be a long period of in-between-ness,
navigating generational structures of care in a society also not sure if they are children or
adults either, and this structural awkwardness will also remodel the patterns of their health care and practices.

Conclusion

Three years on from the day I sat with children arguing over my lunch, a new organisation initiated a campaign calling for the government to supply all New Zealand school children with lunch. The Eat Right, Be Bright campaign (www.eatrightbebright.org.nz) draws on the language of children’s rights to argue against ideologies of parental responsibility, while pointing to successful programmes overseas which show that uptake of school lunches is high, stigma reduced, and health benefits enjoyed by children of all economic backgrounds when nutritious, appealing meals are supplied to everyone. A concurrent media article (Nikula 2018) calculated an annual cost of $720 million based on the well-established and comprehensive Finnish model. Still, the New Zealand government has rejected far more modest proposals targeting primary-school aged children at low decile schools which budget between $3.4–14 million per year.

State policy on this issue, therefore, is not so much a matter of economics as it is of ideology. The services supplying provisions and health care to Tūrama School are, directly or indirectly, the products of neoliberal state views on responsibility for children’s care, whether that be a free fruit programme wrapped in health promotion, or corporations capitalising on the vacuum created by the state’s abdication. In passing off this issue to NGOs and corporations, however, the state also vacates the possibility of care that is child-centred, coordinated, and adequately resourced to alleviate the issues arising from poverty. KidsCan does not have the funds or infrastructure to offer an appealing and universal lunch, while Fonterra has little motivation to supply a non-dairy substitute for lactose intolerant children who may be equally in need of calories and protein. The rheumatic fever clinic, operating under a small and specific community contract, can only refer families for housing support under tight criteria, while accountability for the structural conditions underlying children’s poverty and ill health remains contested.

The nature of these services, though, is not wholly determined by adults, because they are interpreted by children who hold their own, different ideas about what responsibility for care means. From children’s perspectives, adult, and society’s care is part of what constitutes childhood. The degree to which these services operate in congruence with children’s
generational understandings of care therefore shape how children bring these services to life and transform them through social activity. Children collectively make spare lunches into stigmatised care, while, together with teachers, they generate ritualised practices of consumption from the milk and fruit that is supplied to everyone. Younger children enjoy their status-affirming care from Whaea Allison, but those transitioning into adolescence experience friction when this care reaffirms their status as children, and consequently form new patterns of engagement. Thus, children make culture within the structures created by society, but these cultural meanings and practices also, in the Geertzian sense, constrain them in their own webs of meaning, structuring each other’s future practice. Children’s care, and subsequent health effects, is a form of coproduction between the neoliberal state and the practices of children who interpret care within generational frameworks.
Chapter Six
Embodying Inequality

Life for many children at Tūrama School involves coping with bodies that never quite seem to have enough of everything they need: food, sleep, warmth. In winter, particularly, I watched bodies shiver or withdraw, yawn and sneeze, the tactical, homeostatic responses of systems trying to compensate for physiological and psychosocial stressors. Yet their conditions—poverty, institutional structures, even the climate—also seemed to restructure children’s bodies in deeper ways, hinting at habituated modes of adjusting to chronically suboptimal environments when there is little else that can be done. Bodies that shiver and emote and feel pain are functioning as they should, but bodies that do not perceive hunger when they have no food, that are not anxious under threat, or that are numb to the cold, have learnt to accommodate enduring conditions they cannot change.

My analysis of these habituated bodies comes at a time in childhood studies where, after a turn towards the socially constructed nature of childhood, the body has re-emerged as a focus of children’s lived experience: both how the body structures children’s social experience (by size, shape, age, gender etc.) and the ways in which the social world structures children’s bodies (civilising, regulating, and disciplining) (Hörschelmann and Colls 2009; James 1993; Mayall 1996; Prout 2000b). Increasingly, this line of childhood research has followed Prout (2000a, 2005) in viewing the body as neither essentialised nor separate from the social, but as the material foundation of children’s social experience and produced in concert with the social. Already central to children’s lives, the body is especially implicated in the study of health and illness, where health care practices are often performed in response to experiences of the body—symptoms, sensations, emotions—and thus rely on ‘reading’ and interpreting bodily signals. Yet these experiences are not universal or standard, but are patterned by local social context through embodied learning and the cementing of practices over time—the forming of habituated bodies.

An understanding of children’s health, therefore, must begin with an understanding of the processes which shape how children may come to experience their bodies differently in different contexts. The literature concerned with embodied childhood has long recognised the
way that schools, in particular, regulate children’s behavioural, cognitive and emotional development (Christensen 1999; Christensen, James 2000; James, and Jenks 2001; Mayall 1996, 1998, 2002, 2015; Prendergast 2000; Prendergast and Forrest 1998;). This research reveals the power structures embedded in the way adults constrain and shape children’s bodies and the role of children’s agency in this negotiation, but tends to stop short of a detailed biosocial analysis, with less attention to the specific processes of the body that are implicated and how these feed back into the social experience. Parallel research in biological anthropology picks up where the social research left off, diving under the skin to trace how different structural environments manifest in distinct biological signatures (e.g. Flinn 2011; Panter-Brick, Todd, and Baker 1996), but tends to position children as passive absorbers of their environment, omitting the interpretive elements of bodily experience that make this a dialectic process.

This chapter aims to bring these two literatures a step closer together, by considering the biosocial production of the experience of the body, with a focus on how children’s internal perceptions of their body are socially calibrated. Using the examples of temperature, hunger, and sore throat, I unpack the processes through which children coproduce their bodies at the intersection of local resource distribution, and both adult and child cultures, in the overlapping fields of home and school. I argue that bodily perceptions are biosocially learned and accommodated, becoming part of the habitual practices of the self, and that these experiences, and their subsequent health care practices, can be implicated in the production of unequal bodies. Children not only learn to interpret their bodily perceptions in culturally mediated ways, but this learnt process in turn structures the perception and experience of the body, shaping bodily practices, and coproducing the biosocial body itself.

The ‘reembodiment’ of childhood

After a major shift in attention towards the socially constructed nature of childhood, the turn of the 20th century saw a ‘reembodiment’ of childhood research coinciding with renewed interest in the body in social theory more generally (Prout 2000a). While previous research focussed on embedded power structures and the significance of the discursive to children’s experience of the body, some authors (Colls and Hörschelmann 2009; Prout 2000a, 2005) have challenged the ‘disembodiment’ of childhood, critiquing the implicit Cartesian dualism reproduced in a focus limited to how bodies are socially produced and experienced. In
particular, Alan Prout has expanded upon Shilling’s notion of the body as ‘unfinished’ in relation to childhood. In his (2000b) edited volume, researchers from social anthropology and sociology took up an approach to embodied childhoods that views the biological and social bodies working in synthesis to model and remodel each other over the course of the lifespan. While children have long been considered incomplete versions of adults, this notion that adult bodies too are unfinished has sparked more serious consideration of the processes through which the social and physical bodies develop dialectically, rather than fixed by distinct stages of biological development (Worthman 1993). This thinking parallels a shift in biological anthropology which likewise saw increased attention to the ways that the social environment becomes embodied in the developmental and microprocesses of the body, shaping trajectories of growth and patterns of morbidity and mortality (e.g. Panter-Brick 1998).

Under this paradigm, researchers working at the nexus of medical anthropology/sociology and childhood have examined how the body is implicated in the lived experience of childhood through the embodiment of social relations; i.e. the way that the body forms the basis for social relations and so shapes and codifies, while in turn is shaped and codified by, the social world. Key markers of the body include age, used particularly by school institutions to designate status (by academic class), and body size, which constitutes standards of normalness and difference both for medical professionals and for children in peer cultures (James 1993, 2000). In this way, the body mediates social experience; children have variable subjective experiences of the social world according to how they look and behave in comparison to their peers, and individuals’ experiences change as their bodies do.

Yet while the social is grounded in the materiality of the body—its size, shape, performance, gender and appearance (James 1993)—the social experience also mediates the body. In addition to shaping the body itself, social expectations and experiences also influence how children learnt to be in their bodies: seen through comportment, behaviour, emotions, sensation and health. A main focus of this literature has been the institutional environment, where health practices are structured according to the agenda of teachers and the formal curriculum, in contrast to the home environment where children have much more autonomy in their health care practices (Mayall 1996). Authors have drawn from Bourdieu’s habitus (1984; 1977) or Elias’ concept of civilising the body (1978, 1982) to theorise the processes through which the school environment initially becomes embodied, and emphasise how these habitual practices linger in the body long after school has ended. This literature examines the
way in which the school regulates, civilises or disciplines bodies (Mayall 2002; Valentine 2009), in addition or contrast to the socialisation that occurs at home, and with the focus on the way embedded power structures enable adults to shape children’s bodies and restrict their bodily autonomy. The institutional environment also uses disciplinary techniques to compel children to subdue or control their bodies, including shaming and punishment. Those better able to habituate their bodies appropriately to fit the school are less likely to be on the receiving end of disciplining tactics or social stigma. For example, Christensen, James and Jenks argue that children’s bodies, disciplined to ‘eat, sleep, wash and excrete, mostly, at specific and regular times’ (2001:208) internalise and become patterned by the temporal structures of the day. This is not a one-sided process however, as children’s agency plays a role in this interaction. While bodily discipline is promoted by teachers who require children to sit still and work quietly, by deliberately adopting the correct bodily posture and appearance, for example by pretending to look for a book off the shelf, or bending the head over a page, children can strategically ‘pass off’ whispering to friends or taking time for oneself as ‘work’. Hence, as much as schools shape bodies, by acting within and responding to the structures of school, children also play a major role in the coproduction of their own bodies.

Such behavioural modifications are often not temporary, but become absorbed into the habitus, and migrate across fields, interpolating the home habitus as well (Mayall 2015). While the influence of schools on children’s bodies can be considered as benign or pastoral (Christensen, James, and Jenks 2001), and certainly it can be useful for a child to regulate their bodies for other contexts, this line of research reveals how school can be taken-for-granted as a powerful mediator of child—and eventually adult—bodies. However, these analyses tend to stop short of a deeper unpacking of the dialectic between the social and the biological processes of the body, which could extend understandings of how the social environment shapes embodiment. Institutions such as schools structure the way that children learn to read and interpret their bodies in quite fundamental ways, and in some circumstances this may contribute to human variation and less-benign disparities between children.

In biological anthropology, research has shown how social conditions interact with biological process to produce differently embodied childhoods. Working with children in the Caribbean, Mark Flinn shows how family structure and events can alter cortisol profiles, which directly contribute to illness events (Flinn 1999; Flinn and England 1995, 1997; Flinn, Ward, and Noone 2005). Similarly, Catherine Panter-Brick shows how the diverse social living
conditions of homeless, rural and urban dwelling children in Nepal impact stress responses (Panter-Brick and Pollard 1999) and growth (Panter-Brick, Todd, and Baker 1996). Such research reveals just how deeply the social environment is woven into the micro-processes of the body. However the quantitative nature of this research tends to collapse the social into broad categories, for example ‘street children’ versus ‘school children’ (Panter-Brick, Todd, and Baker 1996). So on the one hand, the embodiment literature from social anthropology and sociology unravels the role of the interpretive and the structural in children’s embodiment, while the biological literature teases out the micro-processes that are biosocially shaped underneath the skin. My hope here is to bring these two literatures a step closer together by extending consideration of bodily processes within an interpretive framework.

This perspective can be seen in the issue of children’s toileting practice, where scholars have described the authority of U.K. schools over children’s toilet access, illustrating how children’s self-care practices at home contrast with their lack of autonomy at school (Mayall 1996, 1998; Prendergast 2000). Invoking a children’s rights discourse, this literature documents toileting at school as a source of discomfort, shame, or distress, especially for children who are still developing this technique of the body; Prendergast (2000) describes children being refused to access the toilet, and the embarrassing situations that could arise when children wet or menstruated through their clothing. Poor sanitary conditions and perceptions of safety and security of school bathrooms can also cause children to avoid going to the toilet at school (Lundblad and Hellström 2005).

However recentering the body shows that the control of schools over children’s toileting practices is more than an issue of rights or emotional wellbeing. Prendergast (2000) unpacks the effects of school on the body in this regard most thoroughly, describing a ‘weighty mindfulness’ of menstruation burdening girls across U.K. schools—a situation where girls must be constantly and completely mindful of this process of the body, having to predict, prepare for, negotiate, and protect their periods, with strong social sanctions and stigmatisation befalling any girl who fails to manage her period unobtrusively. However, from here questions emerge about the processes through which the biological body and the social become intertwined: how does this affect the way girls learn to ‘be’ in their bodies, their perception of the signs of impending menstruation and possibly the subsequent construction of pre-menstrual syndrome? Likewise, for children in schools where toileting is

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1 Emily Martin asks similar questions of how medical perspectives and work structures shape women’s experiences of menstruation (1999, 1988).
closely controlled, quite complex bodily processes—taken for granted by adults—are asked of them. Children are required to predict when they might need to go to the toilet in future, and go to the toilet before the urge is strongly felt. This requires a ‘tuning in’ of attention to the signs of particular parts of the body, in order to remember to check in with and notice subtle sensations indicating the state of one’s bladder, at particular times or in particular spaces. This represents a way in which children’s bodies structure the social—shaping teacher and children’s practice—but also of how the social shapes the processes of the body. As children learn to become attuned to these particular signals of the body, they develop a practice of perceiving the body: a synthesis of the body’s biological signals structured by the social (in this case temporally and spatially) which become cemented into the physical self.

Toileting restrictions may also have health effects; evidence suggests that limiting access to the bathroom may alter bladder and sphincter function and is considered a factor mediating the development or persistence of urinary tract infections and other issues (Cooper et al. 2003). Thus, a closer focus on the particularities of body-social relationships shows how the institutional world of the school shapes children’s bodies in very fundamental ways, in the internal calibration of attunement to and interpretation of bodily signals which may have further implications for health, as well as psychosocial wellbeing. My aim in this chapter therefore is to push the analysis of the body further within this literature, via more detailed examination of the biosocial microprocesses that solidify into variation in ways of being in the body which can germinate variable patterns of health.

Bourdieu and the body

Regulation, body work and other such practices of the body, shaped by local concepts of what bodies should be in a given context, solidify over time into the habitus, meaning that children who have experienced similar social conditions embody these in similar ways. As described in chapter one, Bourdieu’s concept of habitus refers to a ‘structuring structure’ (Bourdieu 1984: 170); the way that people come to embody their social environment through routinised participation in structured social systems which predispose them to perceive, think and feel— or ‘taste’—in culturally guided ways. Thus bodily practices are the reproduction

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2 Similar processes may occur in domestic spaces, such as toileting before bed, but to a lesser degree than at school, and these routines may be more tailored to the needs of individual children at home.
(and reinforcement) of those culturally patterned tastes which then shape the way the world is perceived.

Such bioculturally constituted practices shape not only perception of the world, but perception of the body itself. A useful concept from phenomenology is Husserl’s (1962) notion of intentionality, or the ‘toward-which’ orientation of our being in the world. As Desjarlais and Throop eloquently put it, ‘central to the temporal and embodied structure of human experience is the existential fact that we are emplaced in a world that always outstrips the expanse of our being’ (2011:90). Given a world that is far greater than our ability to experience it, we can only focus on limited aspects; a conditioning of experiential focus that calcifies into a culturally patterned habitus. Likewise, even the body itself, at any given time, provides far greater an expanse of internal information than our attention can hold. We notice different aspects of our bodies at different times, and as we tune in to one part—aching shoulders, thirsty throat, anxious breathing—other sensations recede into the background. This is what Csordas terms ‘somatic modes of attention’ or ‘culturally elaborated ways of attending to and with one’s body in surroundings that include the embodied presence of others’ (Csordas 1993:138). As illustrated by the example of toileting in schools above, which aspect we ‘tune into’, and when, and how frequently, is at least partially socially ordered. Children are socialised to tune into their toileting needs according to temporal and spatial cues (break times and spaces). We are more likely to notice hunger at socially designated mealtimes, we tune into pain when describing it to a doctor, and we can have attention diverted away from the body entirely during captivating social events.

Importantly, the social ordering of the body shapes the interpretation of and response to those perceptions, with implications for health: hunger perception helps to shape nutritional intake; temperature perception protects the body from homeostatic stress; and pain perception has a range of health implications, including for social and medical aspects of health care. Moreover, as Bourdieu notes, the habitus does not only organise practices and perception, but is also a ‘structured structure’ (1984:170) where distinctive modes of being are internalised and reproduced within social classes. The examples I explore in this chapter are particular to a social group whose experiences are typified by poverty and social marginalisation, and whose sensory modes therefore represent the perpetuation of an embodiment of social inequality. Finally, the structuring of the senses that occurs in early life may set trajectories of bodily experience and response that continue to adulthood long after the childhood environment has changed.
I emphasise the generative capacity of Bourdieu’s concept, taking habitus as an open and malleable structuring structure that guides, or disposes, but does not determine the course of biosocial development, and which shifts and mutates and opens up new gaps for subcultures to emerge. As illustrated in the previous chapter, children’s peer cultures are important here; children produce their own field of structures which are distinct from adults, but operating at a higher gear of revolution, rapidly mutating to evolve new structural forms as they acquire new information and experience. Often overlooked as a serious force structuring children’s embodiment, as this chapter will show, the intersection of children’s cultures with institutional structures and resource constraints can shape a suite of practices which then become embodied—in this case, through a modelling of perception.

The body as barometer

A challenge of bodily ethnography is in knowing another’s experience of the body, when that experience can only be understood in reference to oneself. As such, a great deal of this chapter is speculative, composed of scraps of evidence that hint at a different experience of the body. I cannot really assess what children are feeling. However, in drawing from my experience of my body as a form of data, I follow in the tradition of sensory ethnographers like Robert Desjarlais and Loic Wacquant who engage in ‘participant sensation’ (as opposed to observation) or ‘feeling along with’ to reveal new perceptory orientations (Howes 2006). Desjarlais (1992) uses his bodily experience to access an understanding of shamanic rituals, while acknowledging the limitations of his ability to truly experience his body in the same way the Nepalese healer to whom he was an apprentice did. Instead, through comparisons of his experience with reported experiences of the healer, Dejarlais conceptualises a hybrid form of embodied experience, recognising that his body and its perceptions are founded in his previous life experience, and those continue to shape his experience of the shamanic world. In his ethnography of boxing in inner-city Chicago, Wacquant (2004) documents the changes that occur in his body in parallel to his observations about the bodies of the men training alongside him. Although his background was quite different from his informants in the gym, the comparison of the changes in his body to the bodies of other boxers at different stages of development—the ontogenesis of rhythm, stamina, and bodily discipline produced by social rules and cultural norms—verified the generalisability of the personal. Through this process of pedantic autopphenomenology in reference to their ‘native’ peers, both authors could triangulate their experiences and gain insight into these specific ways of being in the body.
However, where researchers working with adults can access detailed descriptions of how others experience their bodies, this becomes more difficult with children, who are less experienced at translating their embodied knowledge into verbal expressions. My methodology here therefore relies heavily on observation—noticing the times when my perceptions of my body seemed out of synchronicity with how the children were experiencing the same environment—and comparison of my experience with the experience of other adults.

It was common for me to use my perceptions in a different way to the children I worked with; through my additional years of schooling I had developed a trained ear for the voice of the teacher, an eye for spelling errors, and an inner sense of when teachers were looking, and when they were not. The school environment catalysed a reawakening of habitus-past, and like riding a bicycle, that sense of ‘how to be’ quickly returned. I was less at ease on the playground, where the physical growth since I was at primary school meant that my body was now out of sync with my internal sense of what I could do. I could remember how it would feel to swing around the bars and turn cartwheels, and felt how to do it in my body, but in the time since I had been an avid practitioner of the jungle gym I had lost strength and flexibility, while the physics had changed as my body matured. I was left with the embodied memory of my ten-year-old self, a knowledge that was no longer applicable in a thirty-year-old body (I learnt this the hard way—falling on my head). Despite some prior experience, I also did not have close to the same sense of rhythmic coordination to swing poi\(^3\) that many girls at Tūrama School had developed. These were all understandable differences between what my body was habituated to, and what the children did with their bodies, products of different developmental stages, experience and cultural activities. However, there were two recurring times when I was particularly disconcerted by an apparent difference in experience; two forms of bodily perception that I hadn’t considered might be so culturally learnt. These involved thermoregulation—particularly feeling the cold—and perceptions of hunger.

‘Their thermostat is not working properly’: Perceiving the cold

A cavalier attitude towards the cold is a cultural feature of Aotearoa, and it is not uncommon to see even adults wearing minimal clothing and footwear in winter. However, in this region of South Auckland, the temperature drops to an average low of 10°C in July, and can fall as

\(^3\) Light balls on the end of strings that are rhythmically swung to accompany waiata
low as 0°C overnight. A micro-climate created by surrounding mountain ranges, in winter mornings, I would often drive down the motorway south, watching the thermometer drop a degree or two as I neared the town where Tūrama School was located. On a Monday in July, I leave a trail of dark footprints across the frosted grass as I walk into school, and wrapped up in a wool scarf, hat, and jacket, I sit down on the hard gymnasium floor for assembly, shivering against the cold surface. ‘It’s so cold!’ I moan.

‘It’s not cold,’ the girls next to me say. Some of them are wearing sweatshirts, but many are in only a t-shirt. I am wearing four layers.

‘Yes it is!’ I argue. ‘I bet it’s one degree.’ I look up the temperature on my phone, and it is 10°C, and I pout, ‘well, it’s still really cold.’

Meanwhile, Mrs Randall sits on a chair overseeing the two lines of children in her class sitting on the floor. ‘Whetu,’ she says, ‘take that jumper [sweatshirt] off. It’s not school uniform.’ The principal is ‘cracking down’ on correct uniform, she says. Whetu complies, and shivers, stretching his t-shirt up to his chin and hunching over. ‘I’m going to get sick’, he mumbles, although his hacking cough suggests he already is.

In general children of Tūrama School tended to be underclothed in winter, and the striking contrast between how scantily dressed they were, and my many layers of clothing, was matched only by the contrast in how much we were bothered by the cold. While I would complain about the cold—in part attempting to make conversation—the children rarely did, and instead, as happened in the gym, would disagree with my assessment of the temperature. However, there were times when the children did feel the cold—Whetu, above, notices the difference when he is made to remove his sweatshirt. Inside the classroom, children would huddle against the weak heaters that lined the wall next to the mat. In another class, boys in t-shirts and shorts would drape themselves in blankets and shuffle around like old men. One class had collected a large number of goods for a second-hand stall for the school fair, and after the event, children were allowed to claim unsold items to take home. The objects of most intense negotiation were four large oil heaters, reserved with names written on post-it notes. At the end of the day I watched the lucky children bending over backwards, hugging their prize to their chest as they walked off home to families who probably couldn’t afford the power bill to run them.

Yet much of the time, I felt like I was on a different planet—one where frost and wind and rain were a misery unless properly buffeted with wool hats and thick socks and a warm
jacket. I—like the teachers—wore sheepskin Ug boots and a knitted scarf, and I was still cold. Yet children dressed in t-shirts and shoes with holes in them shrugged when I said it was cold. It was not that they didn’t ever feel cold; it was that their threshold for intolerable cold seemed a lot higher than mine.

I was not alone in the sense that I was experiencing a different environment. Mrs Charles, too, described her own sense that the children’s perception of their body temperature was underdeveloped when she found herself instructing children to remove clothing because they were too hot. She tells me:

‘Mila today was wearing her sweatshirt, which wasn’t a school one, and she was wearing her black jacket at lunch time, and she was sweating, and she had a sore head, and she had sore legs. And I said “take your jacket off and take your sweatshirt off and go and have a drink.” And she did all that and then she came back and said “I feel much better.”’

Mrs Charles continues:

‘I didn’t think she needed her sweatshirt on today because it was hot, but she just, “oh I’m cold.” I said, “but it’s not cold.” I said, “here I am walking around…” [in a sleeveless shirt] and yeah she was hot and she was sweating…. So I think that’s got a little bit to do with it. Their thermostat is not working properly.’

Meanwhile, parental discourse about their children’s illness was saturated with descriptions of the struggle to keep their children warm. Mothers battled on two fronts, fighting not only to keep cold weather from infiltrating poor housing, but also fighting their children to keep them sufficiently clothed. Te Paea attributed ten-year-old Tupono’s three bouts of strep throat to the cold winter, describing how the two of them would sleep in their living room ‘because it’s easier to cordon off and keep warm’ but that Tupono would throw off his clothes when he warmed up, ‘then he’ll get up and go to the toilet and he’ll get a chill.’

Adrienne, having finally got the Housing New Zealand house she needed for her family, described the efforts she made to keep them warm in a house that was still ‘frickin’ freezing like hell, like I go through a lot of power aye, ‘cause I’m trying to keep the house warm for them.’

‘And then they wear a singlet’ I laugh, gesturing at bare-armed Victor. Likewise, Aranui described her exasperated attempts to keep her ten-year-old son Wiremu warm to prevent him getting sick with asthma:
‘It would be the no clothes, playing outside with his sister, yeah so they’d both be out in this cold air. And I’d be walking round going put some clothes on like I told you, stay inside but I have to keep them physically inside.’

Aranui also found herself up at three or four in the morning piling blankets on her children who were coughing and ‘curling up’ but had not realised they were cold.

It seemed that I was not alone in finding I noticed the cold—or the heat—while children did not. Instead, children in general were still developing their ability to perceive and respond to environmental or thermoregulatory signals, while parents actively compensated for and socialised their children to notice the cold and stay indoors or dress more warmly. Teachers also stepped into this role on occasion, but because children at school generally did not have additional clothing to put on, teachers tended not to say anything when children were under-clothed. However, Mrs Charles, who claimed to take on a ‘mother hen’ role as a teacher, described in an interview with me how she would tell children to warm up or take clothes off to cool down, explaining:

‘A lot of my kids will have to be told to take their sweatshirts off, take their jackets off. They have no idea of just doing it because their body is being regulated and things temperature wise… And like I said before, I can’t teach them anything. They are not able to learn if their body temperature is not right and they are hungry. Quite often I tell them go and put a sweatshirt on or go and sit by the heater. I don’t mind if they are wearing a pink sweatshirt to school, just make sure [the principal or deputy principal] don’t see it.’

Described here, therefore, is a process of drawing children’s attention to their bodies and instructing them on how to regulate their temperature: a socialisation of bodily perception. This implies that children do not innately know how to notice their discomfort, or recognise their body temperature as the source of this discomfort, or understand the correlation with the environmental conditions in the same way as adults. Instead, children must learn this as part of a wider process of learning to be in their body. With a deluge of sensory information coming from both inside and outside of the body, far more than can possibly be perceived at one time, children learn which signals to tune into, and which to tune out, in ways that are culturally patterned. In Aotearoa, where attitudes towards the cold can be more relaxed and children may be socialised (by peers as well as adults) towards ‘toughness’, the patterning of children’s experience of body temperature may already look quite different to children in other places. At Tūrama School, the convergence of institutional regulations and poverty create another permutation in this patterning which reinforces the structuring of bodily perception.
Constraints on warm: tuning out the cold

The way that the parents draw children’s attention to the temperature of the environment and related temperature of the body is an example of classic socialisation of how to be in a body, similar to, for example, Geurts (2003) description of Anlo-Ewe socialisation towards balance. However, in the example of children at Tūrama School, I argue that children’s learned perception of their bodies is not only shaped by adult socialisation. While children are still learning to notice the cold, in this environment their lack of power and submissive status to adult authority constrains their ability to respond, even if they were tuned into these bodily or environmental signals. I suggest that when children do not have the capacity to respond to perceptions of cold, or hunger, then the sensitivity of their attunement towards these perceptions may become further muted as a form of coping. It is not that children do not feel cold or hungry, but these do not become the focus of their perception; instead their attention becomes habitually directed elsewhere.

In the case of Tūrama School, a uniform policy requires children to wear only prescribed uniform items. Because children often only owned one school jumper or jacket, if this was mislaid, forgotten, or in the wash, many children would come to school underdressed. The uniform policy is not consistently applied—many teachers turn a blind eye to most instances of children wearing non-regulation uniform—however children will often be admonished or ‘told off’ by senior staff if spotted. This unpredictability of adult sanction means that many children will wear a non-regulation jumper to and from school and in the classroom, but remove it to go outside during break times rather than risk punishment. Children could request to borrow an item of ‘spare uniform’ but they tended not to do this out of shyness and fear that they would be embarrassed or questioned.

So, for example, when one chilly morning I sit outside the deck eating morning tea with Marielle, I am concerned to see her wearing only a t-shirt, with a jacket around her waist. ‘Aren’t you cold!’ I exclaim. ‘Why don’t you put your jacket on?’ Marielle shrugs. ‘Nah, I’m not cold.’ She starts talking about something else. But a few minutes later she is huddled against my side, and I see her arms are goose-pimpled. ‘You are cold! Put on your jacket!’ She shakes her head and mumbles that she’ll get in trouble, and I realise the jacket that I mistook for school regulation actually was not.
Children’s ability to respond to feeling cold, therefore, is dependent on their ability to conform to the regulations of the institution in which they are placed. The school also constrains children’s ability to thermoregulate through other methods; in both of these examples children were placed in situations where they were compelled to sit still—Whetu on a cold gym floor for assembly, Marielle on the steps for lunch eating time—which prevented warming through physical activity. The perception of temperature, therefore, is likely developed not only through adult socialisation, but also in relation to distribution of power and authority which creates constraints on access to the material or bodily resources needed to maintain homeostasis.

Yet, while they were not allowed to warm up, both children in these examples were clearly aware of the cold, as I was—although I was the one drawing Marielle’s attention to her goose-pimpled skin while she did her best to ignore it. Meanwhile others, similarly underdressed, seemed unperturbed.

Such variation in children’s cold-sensing abilities is not unexpected given that children’s backgrounds, experiences and degree of development is far from homogenous. However, it may be important to note that in these examples both Whetu and Marielle had previously
been wearing clothing which they were then required to remove. For children who lose
clothing, the sharp change in temperature is perhaps felt more keenly than for children who
had been underdressed for the whole day. Children who have learnt to access what they need
to keep them warm—like myself—have also likely better learnt how to read the signals that
tell them when more clothing is needed. Unfortunately for children who are powerless to
respond to feeling cold, a well-developed sensitivity to cold becomes an impediment to
functioning in other areas, directing children’s attention towards their ongoing discomfort at
the expense of attention to other social or developmental needs. In this context, tuning out
cold enables children to focus more attention on socialising and play; the discomfort of cold
recedes, though likely still inhibiting full functionality. For a child at Tūrama School, an
insensitivity to cold may be an advantage, when there is not anything they could do about it
anyway. However, rendering children helpless about feeling cold potentially also disrupts
their ability to learn when and how to respond to homeostatic strains on the body; what
Freund (1982) describes as ‘being in touch’, or the capacity to monitor and interpret
messages of the body and mobilise resources to deal with those messages. Such a disruption
of bodily signals and their interpretation can also occur in the regulation of other aspects of
the body, including hunger.

The coproduction of hunger

This intersection between socialisation and material constraints as forces shaping children’s
bodily perceptions also manifests in experiences of hunger. Bourdieu (1984) uses the notion
of ‘taste’ to explain the way in which people develop preferences for what is available to
them, where ‘individuals appropriate as voluntary choices and preferences, lifestyles which
are actually rooted in material constraints’ (Shilling 1993:129), and thus the social
environment becomes deeply embodied. Bourdieu discusses this manifestation of habitus in
terms of classed dietary preferences, stout versus champagne, but here I apply this notion
more broadly as a way of understanding the way constraints are given socially constructed
meanings and embodied by the children at Tūrama School. Here, a lack of lunch is reframed
as ‘not hungry’ and this becomes an embodied ‘choice’, which I argue helps to calibrate the
perception of hunger. This modelling of the senses, again, occurs not through traditional
socialisation, but is coproduced out of children’s own cultural rules, which are in turn
influenced by material constraints.
Research across disciplines suggests that signals such as hunger or satiation are intrinsically biosocial, and the social and cultural milieu may variably interpolate the experience of hunger at the moment of noticing, recognising, interpreting, or responding to perceptions. In anthropology, Hastrup points out that ‘while the need for nutrition is universal, the ‘feeling of hunger’ is culturally mediated’ (1993:731); hunger signals can mean different things in different cultures. For example, Schepers-Hughes’ (1993) analysis of ‘nervoso’ among shanty-town dwellers in Northeast Brazil demonstrates a collective reinterpreting of chronic hunger as illness in a context where people had greater ability to obtain medical treatment than sufficient food. An elastic folk illness category ubiquitous across many cultures particularly in the Mediterranean and Latin America, in Brazil a particular medicalised version of nervoso had come to represent the symptoms of collective starvation—wasting, madness, shakiness, irritability, despair—as a personal problem requiring treatment with medication. In this context of chronic food insufficiency and routinised suffering, talk of hunger had become disallowed, fainting or rage represented as a personal deficiency, weakness or nerves. This reflects a shift in the way the body is perceived in relation to its needs and what those needs imply about the society in which that body resides. While a hungry body needs food, a sick body could be treated with medication, and a prescription for tranquilisers and sleeping pills was obtainable while regular, full meals were not. A hungry body indicates a malfunctioning society, while a sick body is the result of personal misfortune. Meanwhile the impoverished of Brazil came to share the hegemonic medicalised view of themselves through a subtle transformation of everyday knowledge and practices of the body. Being prescribed medication to treat ‘illness’ which is clearly connected to malnutrition reinforced that people and their children were indeed suffering from sickness, a distortion of reality where the hungry become complicit in reproducing the ideologies and practices that deny what they really need.

In this way, hunger signals may be culturally interpreted or reconstructed as something else, but cultural and social norms can also shape whether or not hunger signals are perceived—or perhaps even produced—in the first place. In psychology, experiments have pointed to a variety of external signals that can cue people to consume food, including sensory cues that are exploited in advertising, and culturally normative cues such as portions on plates and structured times of day for eating (Herman and Polivy 2008; Wansink, Painter, and North 2005). For example, Rozin et al. (1998) conducted an experiment which found that amnesiac patients who had no recollection of just having
consumed a meal would consume a second, and then begin a third lunch when the meal was presented to them 10-30 minutes after the first—despite the fact that their stomachs were physically full. The researchers concluded that the memory of the last culturally defined complete meal was a key factor in mediating bodily cues.

Adult socialisation is also a powerful mediator of children’s eating. While children are born with the ability to recognise hunger and satiation (Fomon et al. 1975), adults can often override children’s own feelings in ways that can disrupt the ability to self-regulate food intake. A U.S. study of adult communication with children at mealtimes reports a variety of ways that adults cue children to modify their eating, including drawing children’s attention to their internal cues—referencing hunger, thirst, or fullness—but also to external cues such as food quantities or the time, encouraging children to eat more, or telling children they are done (Ramsay et al. 2010). In a seminal experiment, Birch et al. (1987) showed that children who are conditioned to notice external cues—for example the amount of food on their plate or rewards for eating—showed less responsiveness to feelings of hunger and satiation than those who were cued to focus on their internal state. The implication is that if the perception of hunger and satiation can be disrupted through social cues, then the relationship between bodily cues, perception and interpretation must be the outcome of a biosocial coproduction (Lock 2001); a learnt process. And if the production, awareness or encoding of bodily signals is biosocial in nature, then it follows that when social conditions differ, the way that bodily signals are learnt will be different too, resulting in different embodiments of things like hunger and cold.

Embodiment of ‘not hungry’

As described in chapter five, at Tūrama School, every child has the option of requesting a ‘spare lunch’ if they do not come with their own. In theory, the availability of these lunches should mean that every child eats a substantial meal at lunchtime. In reality, the vast majority of children do not. While all children must sit and eat together during the first ten minutes of morning tea and lunchtime—‘lunch-eating time’—perhaps a quarter of them will have a sandwich or other substantial food item (such as a pie bought from the local bakery, now cold), about half will be eating only a packet of chips or some biscuits, and about a quarter will eat nothing at all.
While it is not unusual for children to not eat lunch in other contexts—teachers at a British school describe a ‘sandwich graveyard’ (Morrison 1996)—variation occurs in the degree to which children’s eating habits are seen as ‘normal’ and which receive adult intervention. In the school where Morrison conducted her research, parents would be notified about ‘persistent non-eaters’ and school rules and supplied lunches were aimed towards making sure children had a ‘proper meal’ while supporting parental choice about their child’s nutrition. At Tūrama School, the equivalent ‘proper meal’ is the rare exception, while teachers, having limited ability to influence what children eat, are usually more concerned with the proper disposal of rubbish. This difference in what is seen as normal or acceptable by adults as well as children’s peer cultures, together with material constraints on what is possible, interact to structure children’s eating habits in particular ways. Following Bourdieu, I argue that these habits solidify into routine practices of the body, structuring perceptions, and ultimately the health and nutritional status of the body in locally specific ways.

For several weeks at the beginning of fieldwork, at lunchtime I sat next to 11-year-old Pikau, a physically slight girl who had immediately claimed me as her friend. While I ate my sandwich, Pikau would never eat, instead hunching over her bony knees on the step while she chattered to me about the happenings of the day. Uncomfortable eating beside someone who was not, I would ask her why she wasn’t eating, and for weeks she would reply that she wasn’t hungry, close off, and change the subject. ‘Not hungry,’ I learnt, was the standard response I would receive from any child with this question, as predictable as the lack of lunch in the first place. It took me a while to realise that my question was not considered socially appropriate by these children who would happily answer almost anything else. When I stopped asking, Pikau eventually volunteered that she didn’t have lunch, explaining ‘my mother only makes lunch for my brother, so she can buy me more clothes and toys,’ which did not clarify things for me.

Yet when children had the opportunity to eat, many of them still did not. I was amused one lunchtime when, after Cassidee told me she wasn’t hungry, Dandre gave her a packet of trail mix, which she accepted. Cassidee then proceeded to pick out and eat all of the chocolate

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4 Teacher responses also varied within Tūrama School and at different times, from a blind eye to wry resignation and sarcasm ‘sit here and eat your highly nutritious morning teas’ to the period of intense monitoring carried out over several weeks by Mrs Charles, described in chapter five.

5 This came other variations; ‘I don’t eat lunch’, ‘I don’t like to eat at school’, but was a very common refrain.
chips, and wandered across the field, sprinkling the rest over the grass. It seemed that when she said she wasn’t hungry, she was being quite genuine.

Coming from an adult assumption that a child without lunch is a hungry child, and a hungry child wishes to eat, I was perturbed by the chronic undereating and baffled that so few would take up the offer of a free lunch. ‘Not hungry’ was a foreign concept to me when I had grown up on three meals a day and would, after a morning of intense activity, be even more hungry than I would be on a non-school day. As discussed in chapter four, some of the rejection of spare lunches is likely due to the stigmatised social meaning attached to spare lunches. However, I suggest that the effects of this intersection of material constraints and children’s own cultural practices further create a peer socialisation and embodiment of ‘not hungry’. Children’s embodiment of not hungry is not only about food scarcity, but also how practices of food distribution and consumption are understood as part of a social order and governed by a complicated set of social rules predicated on a sharing economy. In brief, this social order involves an economy of sharing when a resource is plentiful as a strategy for buffering against future uncertainty. While some foods are categorised as ‘sharing foods’, coming with a moral expectation to share, this contrasts with social sanctions shaming against begging for food or ‘scabbing’. As an alternative to scabbing, children may instead use aggressive techniques to intimidate and recategorise food as ‘sharing food’. For those less able to reciprocate in the sharing economy because they rarely or never have lunch, the practice of ‘not hungry’ becomes an alternative to ‘scabbing’, incurring a social debt, or being seen as pōhara, or poor (see chapter five).

At Tūrama School, as in other places, adults attempt to discourage children from sharing food and drink in order to limit the spread of illness, in particular the streptococcal bacteria that causes strep throat. Many children used the concept of germs to explain the origin of sickness and attributed the cause of specific sickness episodes to the sharing of food or drink. However, sharing is an important aspect of children’s sociality and cooperation, as well as a common method of increasing social capital. Bourdieu’s (1986) forms of capital, where accumulated labour takes the form of social or cultural, as well as economic assets, is useful for understanding the micro-economy that underpinned interactions at Tūrama School. While other studies have emphasised the function of ritualised sharing or trade in expressing and regulating children’s social relationships (Katriel 1987, 1988; Mishler 1979), the scarcity of resources at Tūrama School means that sharing, steeped in a moral code, also forms the basis of children’s economies; an important way of building social capital which can later be
translated into the reciprocal use of resources in the future—similar to the strategies Stack (1974) describes as used by Black kin and community networks in the U.S. Midwest.

A quotidian example can be seen in the circulation of stationary items such as erasers or sharpeners—a micro-economy I participated in and to some extent took advantage of as a way to compensate for my lack of cultural capital in other areas. Important to note is that such stationary items are both essential and scarce; children need them to complete their work, but they are easily lost and children have limited ability to replace them. Consequently, children rely on borrowing erasers and suchlike from peers, and to decline is seen as uncooperative or selfish. On the other hand, an eraser is an asset through which the owner can accumulate considerable social capital, building an image of generosity and helpfulness, which can be traded on in future interactions. However, the more erasers are lent out, the quicker they are lost. So when stationary items are incorporated into the sharing economy, ownership of that item—and the opportunity to build social capital—becomes temporary. When I began participating in the classroom I learnt that the best way to engage with and observe children was to join them in their work, and brought my own case of pencils, erasers, sharpeners and coloured pencils for my own use. These items quickly facilitated my entry into the stationary economy—they were admired, borrowed, and inevitably began to disappear—and then I became the one who needed to borrow things. However, since I could much more easily refill my pencil case than my child-peers—even get reimbursed for my ‘fieldwork expenses’—I held an advantaged position within this stationary economy—never quite one of them. The principle is that those who have excess resources are expected to share—but also gain a lot of credit through sharing, which buffers against the times when resources are scarce.

The sharing of food operates in a similar way, although the rules are complicated by the consumable nature of food; while ten people can use one eraser multiple times, a biscuit can only be eaten once. Therefore, children may be expected and pressured to share perceived excess food, especially if the food comes in a form that lends itself easily to sharing, like a large packet of biscuits or chips. For example, one day I noticed a trail of children eating wafer sticks, and followed them to find Victor, who was offering a large container to the swarms around him while at the same time protesting ‘this is my only lunch!’ In this case, it was not the overall quantity of food—the biscuits were very light and even a whole tin would not have constituted a substantial lunch for a large boy—but the culturally defined nature of the food. A family-sized container of biscuits falls under the category of ‘sharing food’.
On the other hand, the practice of ‘scabbing’ or begging for food from peers is socially unacceptable and stigmatised; annoying for the target, it also represents a shameful public acknowledgement that the child does not have the food they need. To avoid being labelled a ‘scabber’, children may pressure or bully others for a share rather than begging. Performed with a bravado that suggests the demand is reasonable, the more aggressive approach works to redefine the food item in question into the category of ‘sharing’ food, including the moral undertone that comes with it; it would be ‘stingy’ not to share.

Sharing therefore represents a socially acceptable way of redistributing food resources, and social pressure to ‘not be stingy’ can function as a way of prompting children to share. If this strategy does not work, however, children may be accused of ‘scabbing’, which is deeply shameful. One lunchtime, after I produce a sandwich and an apple for my lunch. Kauri, a bigger boy who is friendly but often takes on the role of class bully, leans over to me.

‘Oi Miss, what’s that?’

‘Why are you calling me Miss?’

‘I mean Julie. What’s that?’

‘It’s a sandwich.’

‘Yeah what kind?’

‘Cheese and jam.’ This causes ripples of surprise among the children nearby.

‘Cheese and jam?’

‘Ew!’

‘Nah, it’s actually really good,’ I assure them.

‘Oh, can I have it Miss?’ Kauri asks.

‘Don’t call her Miss, ow⁶’

I smile, but shake my head.

‘Oh please Julie? Please can I have it? Half?’ I shake my head again and start eating.

‘Can I have your apple, Julie?’ Ruby, on my other side asks.

‘No I want it!’ Says Ngapaea. ‘Give it to me.’

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⁶ ‘Ow’ is a vernacular form, loosely meaning ‘you’, associated with Polynesian youth and popularised by the catchphrase ‘not even ow’ from the 2004-2009 animated series bro’Town.
'You guys have got a whole basket of fruit in your class!' I say. ‘I saw it before, it’s got mandarins and bananas.’

‘It doesn’t have apples,’ they complain.

Kauri continues, ‘Oh can I just have that bit of sandwich?’

‘Stop scabbing ow!’ I say, and he laughs, amused at my lingo, but withdraws. At this point I start to feel sorry for him, so I call him back, break the sandwich in half, and give it to him.

Then Ngapaea snatches the half left in my hand. ‘Ew,’ she says, looking at it.

‘Oi! You better eat that now,’ I say, annoyed. She tastes it and makes a face.

‘What is this! I thought it was cheese and mayo.’

‘It’s cheese and jam,’ I say. ‘You stole my sandwich and now you’re complaining.’

‘You should have cheese and mayo, and white bread!’ she informs me. She stuffs the rest of the sandwich into her mouth, making faces. ‘This is the worst thing I ever ate,’ she mumbles through a mouth full of bread.

While I tended to be an easier mark for pressured ‘sharing’, in calling out Kauri’s behaviour as ‘scabbing’ I had tapped into a much more powerful social mechanism than I had realised, shaming him in front of his peers. When I called him back, I recast the interaction into the category of ‘sharing’; a shift that was aggressively taken advantage of by Ngapaea, who did not give me a chance to accuse her of scabbing also, but stole, rather than begged for my food. This illustrates that although sharing is socially promoted among children, and pressuring others to share can be an effective way of gaining resources, it also runs the risk of backfiring—especially if used too often—and the child who asks for food may be cast as that most stigmatised being, the scabber.

The difference between children trying their luck, like Kauri, and those who maintain the practice of ‘not hungry’ is, I suggest, in the degree of children’s ability to reciprocate in the sharing economy. For many children, lunch is unreliable, and they may have something to eat on some days, but not on others. The office lady, in charge of distributing the spare lunches, noticed an increase in demand leading up to the day that weekly government benefits are deposited. Other children were less likely to get lunch towards the end of their parents’ pay-week. Children could also forget, lose, or have their lunch stolen. So for some children, participating in a sharing economy was a strategy to buffer against leaner times. However, for many children, lunches were not just unreliable, but reliably absent. A child who never has
lunch may hope for handouts, or scab or bully food from others at the risk of social acceptance, but an alternative to asking for food from others is to not need food at all.

By ‘not hungry’ I am not suggesting that these children really are not hungry, or that they are not undernourished or not in need of food. Nor am I suggesting that this is ‘just’ their socially accepted discourse. Rather, I suggest this as an example of how the lines between the body and its social performance are not discretely drawn. The ‘not hungry’ body and ‘not hungry’ discourse are not reflections of one another, but mutually constitute each other. In this context, children do not become habituated to eating at lunch time—or de-habituate themselves (as it is more common for younger children to have lunch which has been prepared by a parent than older children). Instead of risking being seen as scabbers, children may cultivate a bodily practice of ‘not hungry’, which in itself conditions those children to be less tuned in to hunger signals, and instead focus attention onto other perceptions; a practice that accrues over time into an embodied habitus of ‘not hungry’. In this way, children’s cultures create the conditions for their own socialised body.

Who has a sore throat?

My impression of how children were experiencing their bodies in relation to temperature and hunger comes from the disconcerting sense of incongruity with how I was experiencing mine. My analysis of how the rheumatic fever programme influences how children are learning to ‘be’ in their bodies is based in part from changes I noticed in the way I related to my body, with the inference that in this case, what I was noticing could parallel how the children habituated to theirs. As I observed teachers asking children about their throats, accompanied children to the clinic, assisted the clinic nurses, and pondered the promotional warnings of the danger of sore throat, I was aware of every scratch and tickle in my own throat. When I came down sick on the last day of term—the cliché of teachers—I spent the holiday convinced that I had strep throat, though without a handy clinic, I never got it checked. It may be a coincidence that during the six-week period in which this chapter was drafted, I twice came down with a cold, the most marked symptom of which was a roaring sore throat. But perhaps not such a coincidence; as I will go on to argue here, there is a vast body of psychological evidence demonstrating a relationship between external cues and subjective symptom experience (Pennebaker 1982).
What I noticed, a ‘tuning in’ to the feeling of the throat, was not a purposeful shaping of children’s relationship to their bodies described in much of the childhood embodiment literature, where parents encouraged or admonished children into a particular way of inhabiting the body. Instead, this internal awareness of the throat came by way of a contingent socialisation; the inadvertent and unseen consequence of a repeated cuing towards sore throat by teachers, parents, clinic staff, the media, and other children (detailed in chapter four). In this way, sore throat becomes highlighted as a significant signal of the body because children’s attention is regularly drawn to it: they are asked to notice if they have a sore throat every day, they are affirmed and treated seriously when they self-identify as having a sore throat, and sore throat is linked to anxiety through media portrayals and children’s own peer discourse of sore throat. The intervention of the rheumatic fever campaign and throat clinic in this way becomes a new ‘structuring structure’, generating a new form of habitus specific to this generation of children and in this particular nexus of ethnic background and class.

This elevation of sore throat within children’s consciousness is reflected in the large numbers flocking to the clinic, a corresponding increase in my own sore throats, and in how often sore throat made an appearance in children’s sickness discourse. In brainstorms of ‘kinds of illness’, an activity I conducted during interviews, sore throat was among the most frequently mentioned symptoms, followed closely by rheumatic fever. Unsurprisingly, when I invited children to tell me about a time they remember when they were sick, many stories involved trips to the doctor or school clinic for sore throats. However sore throats also made appearances in unexpected places. When I interviewed children specifically about their asthma, on a number of occasions children made reference to sore throat. For example, Te Kapua told me the worst thing about asthma is that ‘it’s sore in my throat’, and in a follow up interview several months later, when I asked if he’d had any other symptoms of asthma that month, he described:

I usually play outside but when I- when I had… a sore throat, and um… and the asthma came up, I couldn’t go outside? So… sore throat?7

7 In general, children’s descriptions often revealed an understanding of asthma based on their experience at the time an adult ‘told’ them they had asthma. As in Young’s sickness model, children did not tend to parse out the same cluster of symptoms as the biomedical model, or even their parents did, and instead asthma was frequently related to ailments like vomiting, itchiness, headache, sore throat or heart problems. While children report that they know they had asthma because a doctor or parent told them so, it is apparent that the adult did not necessarily explain which aspect of the child’s illness is the ‘asthma’. ‘Asthma’ therefore became associated with whatever children were experiencing during that illness episode.
As a general symptom of childhood illness, mentions of sore throats would not be unexpected among groups of children who did not have a dedicated ‘sore throat clinic’. By comparison however, in another New Zealand study of children’s illness experience, this time with four-year-olds, sore throat is only mentioned once (McIntosh 2013). Also notable is that although the Tūrama School clinic also carried out skin checks and treatments—in fact 130 assessments over the period of my fieldwork—this was a much less emphasised aspect of the clinic, and, though a common occurrence, children rarely mentioned skin infections. Among countless everyday physical and social ills—colds, eczema, asthma, undernutrition, tiredness—sore throat emerged as culturally salient; unsurprising given its social highlighting from the adult world, but reinforced through children’s shared, visible experiences. That such social highlighting should have an effect on actual perceptions of sore throat is consistent with what a large body of psychological literature would predict.

Psychology of symptom perception and its cultural elaboration

The sore-throat clinic intervention—asking children everyday if they have a sore throat—emerges from positivist assumptions of a sore throat as an objective symptom like bleeding or fever: that if a child has a sore throat, they will already be aware of it, and if they don’t have a sore throat, they will know. Of course, like hunger and cold and other forms of pain, sore throat is a subjective symptom, and—if children share the adult understanding of the term in the first place—one that is produced out of a complicated set of socio-psycho-biological processes. Given the vast psychological literature documenting the relationship between external cues and subjective symptoms (Pennebaker 1982, 2000), it seems likely that children’s experience of sore throat at Tūrama School represents field evidence of at least some phenomena that reliably occur in experimental settings.

Of particular relevance is research which demonstrates how attention to a given sensation can alter how people can perceive sensory stimulation. In a classic series of studies, Pennebaker and colleagues investigated the effects of ‘priming’ on symptom perception, where participants who are primed to expect or notice symptoms did indeed report the symptoms as expected compared to control groups (Pennebaker 1982). This ‘priming’ may be seen as analogous to the prompting of children to notice sore throats at Tūrama School. In one experiment, after filling out a checklist of common physical symptoms, Pennebaker’s participants were asked to close their eyes and concentrate according to the one of four
attention conditions: noticing nasal congestion (congestion increase), free breathing through the nose (congestion decrease), sensations that occurred while breathing through the nose (congestion neutral), and a non-bodily distraction (n=48). After a second questionnaire, it was found that groups’ perceptions of congestion changed significantly according to how they had been directed to attend—those who were asked to notice congestion reported increases in congestion; those who noticed free breathing reported decreased congestion—while neutral and distraction subjects changed very little. The effect was so great that three participants blew their nose directly after the experiment—all in the congestion increase group—while a participant from the decrease group reported her nose had never felt as clear in weeks, and suspected the room had been sprayed with a decongestant. It can be imagined how a similar effect may occur when children are asked every day to notice whether they have a sore throat.

Such evidence of psychological mediators in symptom perception offers insight into the way in which a shared cultural environment could shape the perceptory experiences of a collective group of people through psychological mechanisms, creating local variations in sensory modes. Such psychological mechanisms are likely not universal, but also shaped by local cultural meanings and values which influence the interpretation of bodily sensations, for example translating symptoms into culturally recognised categories of ‘sickness’ (Young 1982). As detailed in chapter four, Christensen (1999) demonstrates that as children learn to interpret and classify their subjective experience of the body, they come to understand firstly that some bodily sensations of discomfort or pain are accepted as ‘normal’, and others are not, and secondly that some sensations indicate the need for intervention or treatment, and others are acceptable. In this case, children at Tūrama School are learning that the symptoms termed ‘sore throat’ are important and threatening, while for children in other areas of the country—white, more affluent children—sore throat appears to be a much less salient aspect of this classification process.

However, I want to emphasise that an important part of the process is learning not only to interpret and classify bodily sensations, but to learn—or unlearn—to notice and feel signals like hunger, cold, and sore throat in the first place. As Husserl describes intentionality, amongst a sea of sensory inputs, perception faces ‘toward’ some as others recede (c.f. Desjarlais and Throop 2011:90), including perceptions from within the body. As one feeling amongst a body-full of various discomforts, pains, and even pleasures, the internal state of the throat is not something that children are necessarily conscious of. Even children suffering from severe cases of strep throat, with swollen ‘soccer balls’ for tonsils, would often be
picked up in class checks because they hadn’t come forward for treatment. Perhaps, Whaea Allison thought, this was out of fear or wariness, or because they just hadn’t realised this wasn’t normal, or because they did not know that this is what was meant by ‘sore throat’. It seemed, however, that sometimes children were simply not aware of any soreness. Like the ‘not hungry’ and ‘not cold’ children, for these children, sore throat was not something they had learned to recognise, and possibly even notice.

On the other hand, once children have learnt to notice a bodily signal, they may become particularly tuned into that sensation, and this may help to explain the large number of children coming to the clinic. The regular discursive ‘highlighting’ of sore throat, the routinised surveillance from clinic staff and from children themselves as their teacher daily draws their attention to the feeling of their throat, and the imbuing of sore throat with anxiety and threats of death, brings a parallel 'tuning in' and heightened awareness—a 'highlighting'—of the feeling of one's throat. For most children who presented regularly, coming to the clinic appeared to be a positive experience of receiving adult care and attention. Such positive reinforcement made it more likely that children who experienced a sore throat would come to be checked again; however it also works in tandem with the provocative promotional campaign to motivate children’s awareness of their throat and the sensitised perception of any small discomforts. As children became more aware of sore throat, they become more sensitised to the subtle variations of sensations of the throat, picking up the tickle of ‘eating scratchy food’, the dryness of thirst, and the tenderness from ‘talking too much’. Without understanding the etiology of strep throat, all of these are classified as the same, threatening ‘sore throat’, and verified as such through the clinic processes; a self-referential feedback loop that reinforces the prevalence of this risk. Moreover, children who do become habituated towards sore throat are more likely to get their strep throat picked up and treated, meaning that variation in habitus becomes magnified by corresponding variation in health care. This marks the beginning of biosocial differentiation that can end in unequal bodies.

Conclusion

This particular confluence of resource limitations, cultural norms, and social interventions collectively lay out the structure for a form of habitus where the state of the throat emerges in prominence while other bodily sensations such as hunger and temperature recede. While
there is variation—not all children experience their body in this way—I suggest the presence of a trend towards this particular configuration, also depending on personal circumstances that shape children’s access to resources.

This represents a way of ‘being’ in the body that is not only culturally distinct, but particular to this subgroup of children at this time and place. Children may develop perceptions of their bodies that may be distinct from adults in the community, shaped by ‘structuring structures’ that vary from the environments of adults, and including structures that children themselves generate. At a structural level, constraints on children’s ability to respond to perceptions of cold or hunger may work to mute those perceptions as children ‘tune out’ of discomfort that they are powerless to alleviate, and ‘tune in’ to other stimuli such as social interactions. Meanwhile, children’s peer cultures comprise complex social norms and rules which shape how children may express or respond to perceptions, potentially influencing the experience of those perceptions. Finally, social interventions—in this case, because of new health policy—may inadvertently create new structures that condition children to experience their bodies in a particular way, enhanced by the social meanings children create around these interventions for themselves. In these ways, children can be seen as coproducing the conditions for their own embodiment. While children’s perceptions of bodily signals may, as they grow up, eventually converge with those of their parents, there is potential for childhood tuning into sore throat or out of hunger to set in place trajectories of bodily practices that are carried into adulthood, with particular implications for health. Turning the lens of analysis close to the body can therefore reveal how deep the biosocial dialectic—and the foundations of the social production of health—play out.
Chapter Seven

Practicing Resilience

At ten years old, Cassidee was close to the shortest in the class, but held herself with a kind of calm poise that always gave me the sense that I was already looking at her adult self, just in miniature. She had mastered the cool tilt of the chin and raised eyebrow that adults would use to greet one’s ‘bro’s’, and while friendly enough, she had an insularity that suggested you could talk to her or not—she would be content either way.

Cassidee had been through five changes of school in as many years, and when I interviewed her in term three she commented—as if it were of little consequence—that her aunt intended for the family to move again at the end of the year. It was common for children who were new to the school to associate more with me. I was a secure base to work from as they tackled the task of negotiating foreign social networks and finding their place in the system.

At the start of the year, Cassidee, new to the school and half the size of some children, hovered at the outskirts of the group of girls at lunch eating time, and was the lone girl in her class to go swimming in the pool at playtime. After the pool closed for the winter she would line up for her turn on the spinners, which did not require a social group to play on. When she wasn’t on the playground I might find her wandering alone or with another straggler, and she would come to me for a hug and a chat. When I spent the day in her class, she would sit next to me on the mat. Whenever she saw me—whether for the first time that day or the tenth—she would greet me in a sing-song ‘hi Jool-lay’, to which I would reply with a matching ‘hi Cassi-day’ and she would smile.

Within a couple of months I spotted Cassidee walking along with another girl, Gemma, with swapped shoes so they are wearing one of each. I commented on this to the boy I was standing with, Caleb, and he confirmed ‘that means they’re B.F.F.’s’ [best friends forever]. This symbolic advertisement of friendship tended to occur when connections were newly forming. However, while a tentative bond may have been forged, Cassidee retained her independence, casually joining in games but less frequently engaging in the physical closeness characteristic of intimate friendships.
Cassidee’s father had died, and she usually lived with her mother, who at 45 was considered by the children to be particularly old, and her thirteen-year-old brother, though she has many other siblings she does not know. As noted in chapter two, whānau are often fluid and shifting, and at the time I interviewed her, Cassidee was living with an aunt while her mother had moved in with an elder daughter to help take care of a new baby.

Cassidee’s story is not that of a typical New Zealand child, but it does include many of the features of a childhood life that are distinctly more common among the children at Tūrama School than in other areas of the country—the consequence of generations of structural disadvantage, as described in chapter two. These features: transience, shifting family circumstances, and loss, create a life that is lived on the edge of certainty, not only for these children who are directly affected by instability and grief, but also for those around them who bear witness to and imagine as their own the experiences of their classmates. Produced through a social ecology shaped by economic, political and cultural forces, this vulnerability is not passively experienced by children like Cassidee, but actively responded to through practices that may become culturally normative ways of being among their peers. These practices, I argue, form the basis of resilience, the processes through which children, with agency, employ strategies in response to perceptions of vulnerability to navigate their circumstances.

As a reframing of risk which focuses on how individuals overcome adversity to thrive, the concept of resilience has proved remarkably popular over the last four decades. Yet exactly what resilience is, how it can be identified, measured, and fostered, are still subjects of wide debate (Barber 2013; Bonanno and Diminich 2013; Masten 2001; Rutter 2013). In particular, one question has been the role of children’s coping in models of resilience. This question recognises the tension between structure and agency that is central to the coproduction of health, but also reflects a second tension between competing constructions of children as vulnerable and in need of protection on the one hand, and as having agency and competence on the other (Ungar 2012). Recently, researchers (Ungar et al. 2008; Ungar 2011; Ungar 2012) have proposed a socio-ecological framework to help to resolve the former tension, and such a framework, I would suggest, also serves to reconcile the latter.

While traditional child-centric approaches to resilience have tended to place responsibility on children for coping, approaches which neglect to incorporate children’s experience and activities into the analysis can erase children’s agency or run the risk of environmental determinism. In a socio-ecological view, resilience is produced through interactions between
children and their environment, recognising the constraints and opportunities produced by specific environments while also making room for what children perceive, experience, and do as significant aspects of those interactions (Ungar 2011). This approach also supports the use of culturally specific lenses for understanding local norms of risk and wellbeing, including the way that children’s views may differ from those of adults.

Such a framework for resilience has been successfully applied in some large-scale research (Eggerman and Panter-Brick 2010; Panter-Brick et al. 2009, 2011; Panter-Brick and Eggerman 2012), but not as yet to many ethnographic studies of children. In this chapter I use a socio-ecological framework to unpack dimensions of children’s vulnerability and resilience: how the local environment contributes to children’s experiences of vulnerability, and how they respond to their perceptions of vulnerability through creative practices to harness resources, protect against risk, and negotiate their position within this ecological context. The ethnographic focus here—on a group of children, over time, in the site where they spend a large proportion of each day—allows insight into characteristics of resilience processes that are less visible in the large-scale quantitative studies more typical of resilience research. While being mindful of Ungar’s (2011) call to decentralise the child from resilience research in favour of ecologies, I aim to show that an understanding of children’s ecologies is incomplete without including children’s activities and perspectives, because, as with their health, children are active in the coproduction of their environment and their experiences shape the way they negotiate their circumstances toward their own care practices. At the same time, the socio-ecological resilience concept here helps to reveal new dimensions of children’s practices: as interconnected with other individuals; as sometimes unrecognised forms of resilience; and as accommodations, where buffering in one domain comes at a cost in another.

These features of resilience practices may be briefly illustrated in Cassidee’s story. While some children cling to social connections for the protections they can offer, Cassidee’s experience is that school connections are temporary, and so she has come to mainly rely on herself. The strategies children use to cope with their vulnerability are not limitless however, but closely bound to the specific ecological contexts which children both inhabit and help to coproduce. These contexts constrain and shape children’s social power, the resources and range of practices that children can draw from, while their practices in turn also feed back into their ecology, often creating new vulnerabilities in the process. Cassidee found a resource in the form of the unthreatening anthropologist hanging out in her classroom. But
time spent with an adult means less time spent with peers, and it was only later in the year, when I was spending more time in other classrooms, that she began to form tentative friendships with others. Thus, practices of resilience are often accommodations; they come at a cost, whether that be because practices are associated with new forms of risk, or that a practice that is successful in one domain comes at a disadvantage in another, or that a practice that benefits one child makes another more vulnerable.

Thus by tracing the specific strategies which children at Tūrama School employ, it becomes clear that the vulnerabilities and practices of individual children are inextricably linked to those of the people around them, and that as children operate within their social ecology, they simultaneously coproduce the conditions that open up or constrain vulnerabilities and resources. This is not to say that children are responsible for their environment—the circumstances children face at Tūrama school are largely a result of social, political and economic processes that are beyond the control of the parents, let alone the children. But it is to suggest that individuals cannot be viewed independently of the others with whom they interact, because their vulnerabilities and practices are interconnected.

Practices, here, are also borne through the body, which plays a dominant role in children’s experience and functions as a tool in practices of resilience. A second thread of my analysis therefore spotlights the body within resilience practices as an important locus through which these processes can be traced. Children’s biological immaturity renders them twice vulnerable; they are physically less powerful, but also their immaturity is often used to justify further constraint of their power and agency in the name of ‘protection’. The body is a medium through which feelings of vulnerability can be expressed, but is also a key resource for children’s practices of resilience: through self-care practices of the body, or through using their physical competence to threaten or defend or their vulnerable appearance to elicit care. These embodied practices of resilience contribute to local, and often gendered norms within children’s subcultures, and create new vulnerabilities for children who find themselves at the receiving end of exclusion, aggression or violence. This focus on what children do, and how vulnerability and resilience is expressed through practices of the body, also allows for an analysis of the accommodations made for resilience.

1 This may also be a much more culturally relevant model of resilience as it moves away from a premise of Western individualism.
2 Parallels between the subjectivities of children and women in this way have been noted by Berry Mayall (1994).
The resilience paradigm and socio-ecological frameworks

In psychology, resilience frameworks seek to avoid a deficit model of the relationship between adverse circumstances and poor outcomes by instead focussing on the protective factors that help young people to mitigate risk and achieve success (Luthar and Cicchetti 2000; Masten 2001; Rutter 1987). Risk, in this context, refers to variables that increase the likelihood of immediate or later psychopathy or ‘negative outcomes’ (Goyos 1997, c.f. Boyden 2005). Despite the optimism of the approach, resilience remains a nebulous concept, variably defined, and with great debate over whether it constitutes a trait, a process, or an outcome, how useful it is, what it is really measuring, and the problems of operational definitions of risk, protective factors, and positive and negative ‘outcomes’.

Traditionally, resilience research has oriented towards quantitative, longitudinal studies, with the aim of identifying the predictors of well-being (variably defined) following adversity (Bonanno and Diminich 2013). This research does not usually attend to children’s perceptions and experiences, but relies on normative (i.e. adult) assessments of what constitutes risk, adversity and outcomes, and tends generalise broad patterns at a population level. So resilience may be measured through ‘outcomes’ such as ‘pro-sociality’, physical growth, or school attendance, while ‘symptoms’ or other manifestations of trauma or distress suggest a ‘lack’ of resilience. This research has contributed significant understanding of what influences children’s well-being, in particular, the importance of at least one supportive relationship with an adult (Boyden 2005). Increasingly, however, the normative operationalising of the resilience concept has been problematised, for example through questions about the generalisability of protective factors across cultures. Likewise, the designation of variables as risks, negative outcomes, or signs of resilience have been shown to reflect normative assumptions of the causes and effects of distress. Contrary to what might be expected, for example, Panter-Brick and Eggerman (2012) find that among their Afghan participants, family conflict, rather than political violence, is the most salient cause of distress. Other scholars have argued that individuals can be resilient and also show symptoms of post-traumatic stress disorder (PTSD) (Yehuda and Flory 2007), and that dropping out of school may be an adaptive social strategy for marginalised young people (Dei 1997). This has led to greater attention to context, with an emphasis on defining ‘outcomes’ according to

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3 Layne, Warren, Waston and Shalev (2007) note that there are as many as eight different definitions of resilience in the trauma literature alone.
local cultural concepts of, for example, ‘the good enough life’ (Panter-Brick and Leckman 2013) or ‘doing well’ (Ungar, Brown, Liebenberg, and Othman 2007).

The emphasis on normative values and perceptions of risk is useful for understanding emic views of resilience, but can shift focus from what children do to cope with adversity. This shift is in part a result of an ongoing tension between agency and structure in resilience research. Early trait and coping based research was critiqued for tasking the individual with the locus of control, essentially holding children responsible for their own survival (Ungar 2012). Meanwhile, models of resilience that focus on ‘protective factors’ in environments can tend to position children and young people in a passive role; they develop resilience because of what adults do, either by providing the conditions to facilitate development and inner strength, or by fostering strong social relationships with the child. Shifting the locus of control from the child onto the social and physical environment reinforces the invisibilising of children’s experience and agency, rendering their perceptions, responses, and strategies as by-products of adult interventions.

The socio-ecological approach goes some way towards resolving this tension. Based on Bronfenbrenner’s (1979) bio-social-ecological systems model of human development, the model—developed most comprehensively by Michael Ungar—conceptualises children as placed within ecologies, navigating risk and negotiating resources according to the possibilities available to them (Ungar et al. 2008; Ungar 2011; Ungar 2012). Resilience, therefore, becomes the process of ‘harnessing’ (Panter-Brick and Leckman 2013) or ‘navigating and negotiating’ resources in culturally specific ways (Ungar 2011). This way of thinking about resilience includes children’s agency, while recognising that their resourcefulness is constrained and shaped by the specifics of their local ecological context.

Recent research has applied this framework in large-scale, culturally specific studies of resilience (Eggerman and Panter-Brick 2010; Panter-Brick and Eggerman 2012; Ungar et al. 2007, 2008). In particular, Ungar led an international mixed-methods comparative study of 89 young people aged 12-23 across 11 countries identified as ‘doing well’ in the face of risk according to local norms (Ungar et al. 2007). For example, 14-year-old Saleem painted a relatively positive picture of his life in a Palestinian refugee camp, describing a close-knit peer network who connected him to a collective cause, feelings of self-efficacy and access to political power, all of which enhanced his sense of identity and cultural adherence. He described small collective acts of resistance—for example throwing stones at the Israeli army—that contributed to his sense of power and control, and attending school as a way to
Contribute to the future of a Palestinian state. It is the specific social and political circumstances that create the pathways for Saleem to find a sense of wellbeing in this way, while young people in other contexts will have qualitatively different patterns to how they approach the social injustices, danger and relationships they experience. This is illustrated in the contrast between Palestinian and Bosnian young people’s experiences of political conflict (Barber 2008). While Palestinian youth tended to endorse the conflict and participate in activism or resistance, Bosnian young people’s experience was of senseless victimhood, with little reason or opportunity to engage in their own defense (Barber 2008). The pathways to resilience that young people navigate therefore depend upon the social locations in which context-specific negotiations and cultural constructions of well-being take place. The point is to consider children’s coping as contextualised within environments with particular opportunities and constraints. In this research, therefore, resilience is envisaged as dynamic processes that are mediated by young people’s culture, context, and resources. This represents a shift from earlier resilience research where ‘shopping lists’ of risk and protective factors are employed to discriminate between ‘resilient’ and ‘non-resilient’ individuals, who are dislocated from any socio-economic or cultural context (Layne et al 2007).

This reframing of resilience invites more detailed ethnographic work to examine the particularities of how children experience vulnerability and negotiate risk and resources towards resilience; the ways contexts open up or constrain pathways, and the way these strategies may set children on particular health trajectories. Usefully, within this ecological framework, it is possible to think of children as practitioners in their own health care, recognising their agency and efficacy without putting responsibility on them for coping. I focus here on practices of resilience, which include negotiation of resources as well as the interpretive process of recognising risk, expressing vulnerability, and employing tactics and strategies (De Certeau 1988) in response to experiences. This is important because children’s vulnerability and their resilience are firstly, shaped by their subjective experience, and secondly, not produced independently of the vulnerabilities and resilience practices of those with whom they share their social worlds. Indeed, as this chapter will show, the ways that children experience vulnerability are influenced by the social ecologies they themselves collectively coproduce—ecologies which also enable and constrain their practices of resilience. Finally, children’s practices, while mitigating some vulnerabilities, may be seen as accommodations in that they create new forms of vulnerability for them or their peers. It is thus useful not only to look at children’s resilience in the context of events, experiences, and
cultural values, but also within networks of individuals who are all actively negotiating vulnerability and resilience.

Children’s feelings of vulnerability in understanding their resilience

The majority of resilience research has not accounted for children’s emic perspectives of their own vulnerability and coping. Yet if resilience is conceptualised as a set of interactions or practices, then how children perceive their environment and experience their vulnerability will shape the form that these practices take. Contrary to its appropriation as an index of risk in models of children’s social and health outcomes, vulnerability is a subjective feeling, the experience of which does not necessary co-vary with statistical or epidemiological models of risk. Nichter (2003:14) describes vulnerability as referring to the actual feeling of susceptibility to illness or misfortune; a state of weakness, fear, or worry. In this sense, children are made vulnerable, but they also can feel vulnerable; an experience that may be related to their socially produced powerlessness, but could vary considerably from adult assumptions of what makes children vulnerable (Boyden and Mann 2005; Christensen 2000). What greatly challenges children—such as navigating shifting school peer networks—is not necessarily where adults perceive children’s vulnerabilities to lie. Meanwhile the concerns of adults about children—such as lower academic achievement caused by transience—do not necessarily form part of children’s experience of vulnerability. Because children use perceptions of vulnerability to guide their practices, an understanding of what makes children feel vulnerable, and how they respond to these feelings, must underpin any understanding of their resilience.

Yet children’s perceptions of risk or feelings of vulnerability rarely appear on the radar in health policy which assumes children as passive recipients of care. For example, at Tūrama School, children have a perception of the risk of rheumatic fever that is much higher than the epidemiological risk. This perception comes from interpretations of their experience; as described in chapter four, over half of the children have been given antibiotics for strep throat at some point since the clinic opened, and many take this to mean they either have, or are a step away from having rheumatic fever. Children observe others getting antibiotics and see susceptibility as common. For some children, just the nurse swabbing them is an indication that they are at risk. This, combined with the media message that rheumatic fever can cause death creates a perception of risk that is much higher than the epidemiological
data would suggest: ‘I didn’t want to die. ‘Cause on the ad, him! He almost died!’

Amberlee, age 10, tells me. While in a given population with high exposure to GAS A strains only 3-6% of individuals will develop rheumatic fever (Carapetis, Currie, and Mathews 2000), the children getting swabbed are unaware that they are most likely not even susceptible. Furthermore, it is rare for a child to die of rheumatic fever; most rheumatic fever-related deaths are in adulthood from the complications of rheumatic heart disease (Wilson 2010). The proximity of death suggested by the media campaign contributes to a perception not only that rheumatic fever is common and easy to contract, but that the consequences are more severe than is the clinical prognosis.

Children’s frequent use of the clinic therefore becomes much more understandable when viewed in terms of their own sense of vulnerability. The clinic represents a powerful, accessible resource in mitigating risk through its promise to detect and treat the sore throats that lead to rheumatic fever. The clinic also works with children’s agency; in most cases children decide when they need to go to the clinic and put their names forward, children are free to see the nurse during break times, and together with their parents children may take an active role in managing their antibiotics. The fact that children are not helpless in the face of their perceived risk may work to reduce their sense of vulnerability, and the clinic thus gets co-opted into a practice of resilience.

Understanding this relationship between children’s experiences, perceptions, agency and resources is important not only for understanding their health care practices, but to understand the impact of a health intervention such as the clinic. As adults have inadvertently created conditions where children perceive they are at risk of rheumatic fever, they have also rendered children—and their parents—partially dependent on the clinic to mitigate their sense of vulnerability. When I asked one of the children’s mothers, Anna, how it would be if the clinic closed she visibly panicked:

‘What could I do as a mum, stand and scream. I would stand wherever- the Beehive [Parliament], if I could afford- I would go up there and I would tell them what for. I would rather have my son come in once a week or, you know, however many times in a week to prevent him getting it than have him get it because we don’t have it so local and so close and it’s so convenient than have him get it. Like me and his dad would definitely lose our jobs because there is no way I am having a sick child in hospital and not being with him. And the stress alone would be too much so I reckon we would both lose our jobs, we would have no money to live on… Don’t tell me they’re shutting it down.’
Having had her son Anton come back positive for strep twice, the threat of rheumatic fever is a frightening reality that Anna takes very seriously. For her, the clinic in the school has come to represent the lynchpin holding together a family struggling on the edge of coping. Probably in part due to Anna’s encouragement, Anton was one of the children who presented at the clinic every couple of weeks to be checked. So long as the clinic remained in the school, it gave the family an accessible resource for managing their vulnerability, and in doing so, practicing resilience. Hence, attention to what makes children and families feel vulnerable in specific contexts is essential for understanding how they ‘negotiate resources’ towards resilience.

Children’s vulnerability and resilience in social networks

In the context of Tūrama School, a significant part of children’s experience of vulnerability is produced against a backdrop of a shifting, unpredictable social landscape, where connections are temporary and the peer networks are continuously reconfigured. This local ecology is a result of wider structural factors, including the legacy of colonialism, a housing crisis and demographic changes, but is also co-constructed by the children themselves, who actively respond to their vulnerabilities through creating and strategically navigating their peer cultures (Corsaro 1992; Spray et al. 2018). Thus children’s vulnerabilities are produced through the interaction of structural forces and children’s own interpretive meaning-making, while their responses can create new forms of vulnerability for themselves and others.

I focus here on children’s peer ecology at school, as this is where I was based, but the processes—or characteristics of processes—that emerge are likely to be relevant in other domains as well. I follow a growing body of research documenting children’s peer groups as important sources of social support (Boyden et al. 2004; Boyden and Mann 2005; Spray et al. 2018), though with the caveat that these social practices may not look like the adult view of what is best for children, and that practices that are protective for one child can be at the detriment of another. The significance of peer networks as a cause of distress for children has tended to be overlooked in resilience research in favour of forms of vulnerability that are more visible to adults; domestic conflict, political violence, environmental catastrophe (but see Boyden et al. (2004) for children’s perceptions of peer abuse as the worst consequence of poverty). In this analysis, I aim to bring to light the complexity of children’s practices within the peer ecology they themselves coproduce.
As is reflected in Cassidee’s story, a high rate of transience is a notable feature of school life, and classes experienced a change in membership of up to 25% over the course of the year. Aside from the impact of transience on educational achievement—a factor the teachers would lament—this revolving door of children created a constantly shifting dynamic in classes. The onus was on the new students to integrate themselves, and for many this was a long, slow process. Cassidee, experienced as she was at transitions, took this in her stride, and self-sufficiency seemed to be her strategy for adaptation. She didn’t reject friendships, but she didn’t actively invest in them to the degree of some of her peers. Other new children sought to build intimacies with a kind of urgency, and in the absence of success elsewhere, girls in particular would cling to me, claiming their spot on the mat next to me and pressing their thighs snug against mine. As the weeks went on and their own connections grew, they would gradually leave me behind and find their own place in the social order.

It was hard for those who arrived, and likewise hard for those who left. Sometimes children had a month or two of notice before the move and would talk about it, naming that they were leaving to friends, teachers, and myself. They did not tend to talk about how they felt or what the move might mean, but the naming to others, over again, put words to the significance of the event and their disturbance. The children hearing in turn would ‘tell others’, who would go back to the leaving child for confirmation and ask ‘are you leaving?’ This ‘naming’ would continue up until the child left. Children would also identify whether the child would or would not be present for upcoming events, mapping out the future landscape. When practicing for a choir performance, one girl made it clear to those around her that she would have left before the event took place.

When notice had been given that a child was leaving, a class would sometimes acknowledge or celebrate their last day. Often, however, children would suddenly disappear. Sometimes there had been a few days’ notice; sometimes children would simply not return after the holidays. Sometimes children would know they were likely leaving, but not know for sure, or when. My first encounter with this is when Ben, a friendly boy who often hung around me at play times, tells me that he ‘thinks’ this might be his last day. ‘Oh really?!’ I say in surprise, and comment that he’ll miss being in the school play, which he has a main role in.

‘Yeah,’ Ben replies. ‘I think I have to go live with my auntie.’

‘You think you have to go?’

‘Yeah. My social worker? She said I have to go and live with my auntie.’
‘Oh wow,’ I respond. ‘How do you feel about that?’

Ben shrugs. ‘I wonder what school I will go to?’

‘Do you like your auntie?’ I ask.

Ben smiles, ‘Yeah.’

Ben indeed did not come back to school the next day, and if I hadn’t happened to stop and chat to him while he hung off the rail outside his classroom, he would have been another unflagged disappearance. As a lead role in the school play, Ben’s leaving was of most inconvenience to the teacher directing, who had to hastily replace him. For me, Ben was the first child with whom I had built a relationship to disappear in this way, and I was shocked at the suddenness of his announcement and the uncertainty he expressed in this final conversation; he didn’t know for sure he was going or when, or what his new life would look like, or what school he would be attending the next day.

Moving was hard for those who left, but also could be hard for those who were left behind. Friendship bonds could be abruptly broken, leaving children unexpectedly vulnerable and without their closest ally. Ruby, who we met in chapter two, had been friends with Eponi since the previous year where, as year fives in a mostly year six class and on the edge of social circles, they had paired up. This year, Ruby had found herself in a new class with Eponi, the only classmate who had transitioned with her, solidifying her bond. In an interview, Ruby, supported by her new friends Ngapaea and Alexandra, describes the loss of Eponi several months prior:

‘It’s hard for me ‘cause my best friend left. And I never got to say goodbye because I had to go to my cousin’s tangi [funeral]…’ she trails off. ‘Eponi, she moved to Wellington I think.’ ‘Because of her Dad,’ Alexandra supplies.

Ruby corrects, ‘No, ‘cause of her-’

‘Grandpa?’ Fills in Ngapaea. The three of them have a tendency to build stories in a sort of collaborative project.

‘And so, did you know she was leaving?’ I wonder.

‘Yeah, she told us, she had to go-’

‘-to a tangi.’

‘So how far in advance did you know that Eponi was leaving?’

Ruby hesitates. ‘Um… I don’t know.’

I clarify, ‘I mean, was it like she told you one day and the next day she was leaving, or…’
‘No she told me one day, and then like- oh she stayed for a week. And then on Friday she left, and that’s when I went away,’ Ruby explains. This shifting social landscape leaves children with low social capital and few strong social connections in a vulnerable position. Secure acceptance into a bigger group therefore is enormously beneficial. When I interview Ruby later in the year, she is now a core member of a tight, exclusive group of five girls. Ruby and her two friends describe how their group has explicit rules and roles and wield threat of exclusion to keep membership and behaviour in line. They had kicked another girl, Soraya, out of the group due to the behaviour of her brother. Likewise, they threatened to exclude another girl, Sarah, when she broke the rules and continued to talk to Soraya.

‘So you guys feel safe in your group?’ I ask.

‘Now,’ Ruby explains, ‘now- ‘cause we got a leader.’

‘Her,’ Alexandra and Ngapaea both point to Ruby.

‘And a co-captain…’ Ruby adds.

‘Me,’ says Alexandra. ‘Oh no-‘

‘You’re the choreographer,’ Ngapaea corrects.

‘You guys are quite structured now,’ I note.

Ruby agrees. ‘Yeah because like our- like we’re tryna keep it together but- I’ve been bossing them around, so I have to stop that.’

‘Yeah but um, [Sarah] kept on talking to Soraya and telling her what we’ve been like saying, and talking about’-

‘Cause we got rules,’ says Ruby.

‘Yeah we got rules. Don’t hang out with the haters. Like, haters don’t hang out with the haters.’

‘And no putting your hair on the side.’

In the face of forces out of her control that took away her friend, Ruby here has done what is in her power: seized leadership of a group (instead of one close friend) and created a kind of group culture that promotes the loyalty of those friends, ensuring a more stable social positioning and buffer against future losses.
Though Ruby might be described as resilient, it is perhaps more useful to note the forms that her practices of resilience take within this context, how she has come to arrive at her current strategy, and the effect of these practices for her and for others. Her practices are both produced out of her specific experiences of vulnerability, but at the same time render others (Soraya and Sarah) more vulnerable. In this way vulnerability and resilience are mutually constituting; vulnerability is produced through the interaction of social conditions and the actions of individuals with agency, while resilience is enacted through agentive practices within the constraints of those same social conditions. In a socio-ecological model, this recognises ecologies as dynamic and shifting networks that are coproduced by the children while they simultaneously jockey to navigate their resources in response to perceptions of vulnerability.

Figure 8. Embodied expressions of vulnerability
Embodied vulnerability and resilience

While children hugged me, and cried to me, and interrogated me, and showed off their skills on the playground, the times when they came to silently stand next to me marked a different kind of need. They might lean against me, or sit with their thigh pressed against mine, but they said nothing, and if I asked questions or commented it felt like I was speaking the wrong language. They would stay by me for a minute or two, and then move on, ready to enter their world again. These were the moments for the things that could not be expressed in words. This, I came to realise, was the language of vulnerability.

Stepping away from ethnocentric definitions and capturing the local cultural norms of risk and resilience invites a particular attention to language; the vocabulary, values, and stories that participants use to construct and express the meaning of experiences. This ‘meaning-making’ has been discussed as core to promoting resilience within constructivist or cultural resilience research (Theron and Theron 2011), which identifies concepts or values expressed in local narratives and ties these to a determination to survive or overcome adversity. For example, Eggerman and Panter-Brick describe adherence to six cultural values as forming the basis of resilience in Afghanistan (2010), while the concept of ‘sumud—a determination to exist through being steadfast and rooted to the land’ was drawn out of analysis of narratives from Palestinian young people (Nguyen-Gillham et al. 2008), and in South Africa a cooperative philosophy called Ubuntu or Botho underpins making sense of poverty (Theron and Theron 2011). These concepts, distilled from interviews with adolescents, young adults, or caregivers, codify vulnerability and resilience into language in a way that I struggled to replicate in my observations of the children of Tūrama School—not because children do not make meaning, but because they did not tend to articulate this meaning verbally. Instead, the experiences of younger people may be better read through embodied expressions of vulnerability and resilience; what they do, how they use their bodies, micro-interactions in the context of their local environments. Such embodied expressions are not unique to children of course, but have been noted by other authors as particularly relevant to phenomenological studies of children. For instance, Das (1989) notes that children traumatised by anti-Sikh riots in Delhi would not offer extended verbal accounts of their feelings, but knowledge and memories were encoded in their bodies and expressed through displaying injuries, miming events, and projecting the dead into ghosts whose voices haunted them at night. Memories and meaning making were not bounded within each individual child, but held and signified collectively; for children who did not speak, other children spoke for
them, naming the spot where a parent had died in a way that mirrors the naming of a loss by the children at Tūrama School. Such interconnected expressions of vulnerability could be similarly traced through the embodied practices of Tūrama School children. The stories in this chapter illustrate vulnerabilities that are expressed through silence and touch, and resilience practiced through bodily power, agency, and autonomy, through the embodiment of sore throat and its vigilant care, the language of aggression, or the shared language of pain.

Mila

At age ten Mila had a quiet maturity that both made her an attractive friend to the girls in her class and endeared her to the teaching staff. Bright and watchful, she would get on with her work while the other children squabbled over stolen erasers and exchanged notes. When I wandered around the school, or stood observing, I would often turn to find her standing next to me, watching too.

Mila lives in a two-bedroom house in, according to her, ‘quite a bad neighbourhood.’ At the time I interview her, she lives with her mother and an adult cousin, and shares her bedroom with three of her brothers aged between 16 and four.

Mila loves her father. He has her name tattooed on his back because she’s his favourite daughter, his ‘good girl’, because, Mila tells me, ‘I listen. I… concentrate. I do my chores. I never… I never run away from him- well, one time… Just one time. And I respect him.’ Mila tells me she feels safe when he’s around. He loves her.

Mila explains that her mother is ‘almost dying’, sick with something the family don’t really understand but which the doctor says can’t be cured. ‘And’, Mila tells me in her interview, ‘she’s always got a sore shoulder, also neck, because of my dad. Yeah he, when he was younger, he used to be a Black Power.’

I wonder how her father’s gang affiliations had impacted her mother’s shoulder and neck. Mila explains to me:

‘Well, there was this one, there was this one day, and there was this one day, um, my mum and my dad were sleeping next to each other, and then my dad got up and he was all angry, ‘cause he wasn’t supposed to stay there, ‘cause he was- he was… trespassing? And then, um, and then my dad went outside and slept in his car. And then my little brother jumped into my mum’s bed. And then my dad came and started punching her in the neck? And then he just left without saying sorry or anything.’
Articulate as she is, Mila is unlikely to use the word ‘vulnerable’ to describe herself. Her stories though, explain her silent appearances by my side. She has had a lifetime of vulnerable experiences. But Mila’s vulnerability is interconnected with her mother’s—her illness, her victimisation by Mila’s father—and this creates the context for Mila’s vulnerability. Mila is dependent on her mother’s care, but also sees herself as essential to the care of her mother. Several of her siblings have moved in with an adult sister, but Mila wants to stay with her mother so ‘I can look after her.’ Mila is frightened for her mother’s safety, but she is also worried that if her father perceives that she is taking on too much of the care for the household, he will take her away from her mother, as has happened twice before. As the children understand, illustrated in chapter five, children are not meant to be responsible for a parent’s care, even if they sometimes need to be. In this way Mila’s child status makes her twice vulnerable; her father’s perception of her vulnerability means if she cares for her mother she may be taken away. Yet if she does not protect her mother, her mother’s illness means that Mila may lose her anyway.

Like many of the other children I interviewed, Mila had internalised the warnings of the rheumatic fever campaign and showed a hyper-vigilance about sore throat. She estimated that she had visited the clinic 12 times that year, not because she experienced a sore throat, but just to check, ‘to watch out for it, just in case I get it’. Her sense of vulnerability was in part constituted by her perception of the risk of rheumatic fever, as conveyed through the frightening advertising. But unlike most others, she did not articulate her fear of sore throat in terms of her own illness or death, but because ‘my mum will worry’.

‘Your mum will worry?’ I ask.

Mila explains, ‘and then, she’ll like, take me to the doctor and everything, and then, I’ll tell her that it’s fine? And she’ll be like, no it ain’t, you’re sick! This ain’t a fine thing to think about. And then, I’ll go, ohhhh. And then, my mum says don’t ohhh yourself ‘cause you know that it’s something to worry about.’

Hearing her mother’s concern about the possibility of her illness, Mila hears that her own vulnerability makes her mother vulnerable. In response, Mila goes to the clinic to get checked.

Mila also protects her mother in other ways. When her mother gets out of bed and groans in pain Mila will tell her to lie back down and does the chore herself. And Mila tries to protect her mother from her father:
'I always watch them, like, every five minutes, I’ll always walk into the room and say hi! Hi! Hi! And then they’ll say, can you please go out, and I’ll be like, okay. And then, I’ll walk out, and then I’ll stand next to the door? And like, listen? And, if they’re yelling I’ll walk into the room and tell them to stop yelling.’

On the day of our interview, Mila had been checked with the rest of her class through the class check that happens twice per term. She tells me, ‘I’m so lucky that the throat lady, she came around? And she came to our class. And she said that I was fine’.

‘Today?’ I clarify. ‘How did you feel when she said you were fine?’

‘I felt really happy, ‘cause I knew that I was looking after myself really well.’

Like many of the children, Mila attributed the origin or prevention of sore throat to her self-care, rather than parental care or social factors such as overcrowding. I asked her about the things she does to look after herself and keep her from getting a sore throat. Mila tells me she drinks lots of water, and makes sure nothing bad happens to her.

‘Like, making sure that nothing bad happens to you like what?’ I ask.

‘Like getting kidnapped, or… or like… getting murdered, or getting killed in my sleep.’

I laugh because I think she is saying preventing these things will help prevent sore throat. But maybe Mila is saying that. In a world surrounded by gang conflict, schoolyard aggression and the domestic assaults of relatives, maybe the threat of violence by illness becomes conflated with the threat of human violence.

Children’s bodies as lenses for vulnerability and resilience

The practices of resilience that Mila has carved out for herself are bodily practices; she uses the presence of her body to diffuse tension between her parents, and takes care of her physical health, drinking water and using the clinic to maintain her physical health to protect herself and her mother and find greater security. Her vulnerability is also embodied in her vigilant attention to her throat, and the quiet moments where she stands with me and watches her world. These embodied practices parallel the more cognitive processes highlighted in cultural research with young adults; these children didn’t express their resilience conceptually or in mantras or concepts but it can be read in other languages nonetheless to shed light on where children feel insecure (Andresen 2014) and how they respond. Attention to embodied practices of resilience can also capture local norms of risk and resilience—norms that may be
distinct in children’s peer cultures from those of adults in the same community. These are culturally and ecologically derived ways of using bodies to protect, defend, and move forward in life.

Embodiment as a ‘lens’ for reading vulnerability and resilience is appropriate given the centrality of the body in children’s lived experience. Children’s experience of their bodies is one of continuous growth and development, where universal developmental trajectories are overlaid with cultural meaning. In the West, an emphasis on the ‘becoming’ child means children’s development is very much at the centre of adult attention. Adults measure children’s growth and keep a running commentary on changes, and hence events such as losing teeth and mastering new physical skills are chronicled by children themselves in recognition of their cultural significance (James 1993). Children’s bodies are also a major component of their vulnerability; smaller, younger, or weaker children hold less physical power than those who are bigger, older or stronger. This biological immaturity is used as the justification for ‘protections’ which become embedded in social organisation and policy in the form of segregation, restraints, and regulation, but often limit children’s agency and reinforce their dependency and lack of power (Mayall 1994; Qvortrup 1994). Becoming bigger, stronger, and more adult-like may be desirable as a pathway to greater autonomy and power, though growth is viewed in tension with the desire to be ‘normal’ as determined by comparison with others in the same age group (James 1993). Being too big, as well as too small, can create additional vulnerability via self-consciousness and teasing or exclusion from peers.

Hence the body is central to children’s experience of vulnerability, and so its development and mastery is key to outmanoeuvring insecurity. However, the forms that children’s bodily practices of resilience take may not match with clinical, or even lay adult perspectives of what resilient functioning should look like. The recent shift towards culturally-relevant conceptions of ‘doing well’ troubles adult notions of what is in the ‘best interests’ of children, and is perhaps best illustrated by a burgeoning literature on the resourcefulness and wellbeing of street children (Mizen and Ofosu-Kusi 2013; Panter-Brick 2002). The spectrum

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4 At Tūrama School, mixed year-group classes and a high prevalence of overweight or obesity may have mitigated this effect as the range of shapes and sizes expanded the range of ‘normal’ within the cohort, though those at either end of the wide spectrum could still be rendered more vulnerable.

5 The importance of physical competence to children was demonstrated to me every time I entered the playground; I would be mobbed by children calling ‘Julie, watch me!’ They would rip my gaze away from one another, competing for my attention and acknowledgement as they performed an array of jumping, twirling, balancing, and swinging skills. Sometimes they would demand that I try performing the trick, and would be politely satisfied when I gave in and demonstrated my inadequacy.
of children’s bodily practices includes a range of aggressive or violent behaviours that may be generally unsanctioned by adults in the community, but some of which may be considered normal or a form of resourcefulness for children themselves. This is what Ungar has termed ‘hidden resilience’, or ‘culturally non-normative substitute adaptations’ (Ungar 2011:8). For example, a study of adolescent aggressive behaviour in urban China found that aggression used by young women in Beijing was normalised and viewed as a functional strategy to cope with culturally embedded gender bias in intimate relationships through maintaining personal coherence and resisting negative stereotypes (Wang and Ho 2007). Both female and male adolescents characterised this aggression, which included hitting, pinching, and kicking, as Ren Xing or ‘willfulness’ rather than violence. The same aggression from males would be considered unacceptable violence, but young men were expected to tolerate or accept aggression from their girlfriends. The authors suggest the roots of this normative characterisation may be traced to China’s one child policy, which has resulted in children growing up as the treasured only child, and Ren Xing being glamourised as the acting out of a spoilt child, and expected gender roles, especially in dating, where women are seen to have a more advantageous position than after marriage. This emic view of aggression situates violence as a normalised bodily practice within young people’s subculture, one that is in part produced from the historical cultural context of the wider community, but also through the socio-cultural context coproduced by urban Chinese young people, particularly through discourses and media representations. A culturally-contextualised approach to processes of children’s resilience may therefore need to consider children’s practices not in terms of adult local cultural norms, but in terms of children’s experiences, perspectives, and interpretations.

Children’s aggression as accommodations for resilience

Though proscribed by the New Zealand school system, for the children at Tūrama School, aggression is both a routine practice of self-protection and self-promotion, and a source of vulnerability for both perpetrators and those exposed. I do not think normalised forms of aggression are uncommon among children, nor specific to this context, although lower social tolerance for aggression in more recent times particularly among the middle class may have led to stricter monitoring in some schools. As described in chapters two and three, while New Zealand law prohibits the use of any physical force towards children, talk of ‘getting a hiding’ was pervasive at Tūrama School, and children’s talk reflected a world where violence was normalised, integrated into play, and even a form of resourcefulness. Like in the example
of Ren Xing, the form this violence takes can often be distinctly gendered. For boys, in particular, aggression formed a functional strategy for navigating the vulnerabilities produced in this social world, while amongst girls, aggression towards others could be antisocial, and some girls established practices of harming themselves instead. In employing such practices to cope with their vulnerability, therefore, children can sometimes place themselves or others into new positions of risk. This does not mean that the tactics are not successful; just that successful coping in one domain of life may be at the expense of another.

I borrow from the human growth literature a way of conceptualising this dimension of children’s practices which acknowledges the realities of human practices of coping. While the term ‘adaptations’ is used to describe a beneficial response to poor environments, the term ‘accommodations’ was proposed as an alternative to draw attention to the costs and benefits of coping (Frisancho 1993). This reframing of survival in terms of ‘trade-offs’ challenged the ‘small but healthy’ view of stunted growth that was used to justify inaction in response to the chronic malnutrition of global populations (Panter-Brick 1998a). If resilience, or thriving despite adversity, represents a parallel to beneficial adaptation of biological systems, then accommodation represents a more accurate conceptualisation of the many social processes that play out in the space between vulnerability and resilience. Practices of ‘navigating and negotiating resources’ while effective, nonetheless often come at a cost. This has been noted by other researchers; what Panter-Brick describes as ‘entrapment’, where cultural values can be a double-edged sword when individuals struggle to conform to ‘what makes an honourable Afghan’ (2012:383), or when they constrain or become maladaptive, for example when individuals cannot draw support from relatives because they cannot afford the expected reciprocal hospitality. Ungar too, notes that behaviours that may be adaptive in the short term can carry long-term consequences, though this is not a major focus of his analysis. Nancy Scheper-Hughes (2008) describes different ‘tactics of resilience’, which she illustrates with ethnographic examples from families in shantytown Brazil and revolutionaries in apartheid South Africa. While some of these might be celebrated as demonstrating the strength of vulnerable people in overcoming adversity, Scheper-Hughes notes other tactics might be seen as problematic, or, as she diplomatically puts it, may ‘offend ‘our’ sensibilities and tastes, shaped by very different subjectivities, notions of value, human worth, and the good life, meaning always, the life that is worth living’ (2008:43). These tactics include normalisation, such as maternal acceptance of a child’s death, tactics of daily
improvisation such as using trickery, cunning, or manipulation, ‘getting away with murder’, and socialisation for toughness.

Hence, the responses of the children at Tūrama School may be viewed as accommodations to their circumstances; the responses of active agents within the structures of their ecological system, but responses that constrain options, produce new forms of vulnerability, or that are simply seen as undesirable by those with more power—namely, adults. This is the case with aggressive or violent behaviour, a protective strategy particularly used by boys in the context of peer relationships, but often accompanied by adult sanction, risk of injury, and a loss of educational opportunities.

Nine-year-old Harrison described for me how he had been ‘the funniest and annoyingest in my class’ as he entertained his classmates with amusing noises, but also how his teacher would have to restrain him when he got angry, like when ‘someone was like trying to be like being a dick to me and that… and I was going to kick him in the head, which I did.’ Harrison was diagnosed with ADHD and anger management problems, suspended for three days, and according to him, eventually expelled. Observing Harrison’s entry into playground life as the new boy at Tūrama School, I could imagine how he had come to settle on this aggressive strategy for survival; slightly built, freckled, and one of only three Pākehā boys in the senior school, his fair skin and light hair immediately marked him as a target for bullying.

One afternoon, while sitting on the mat, he showed me the injuries he’d acquired earlier at lunch, deep scratches on his arms and a scrape where he’d fallen down. He was aroused, talking quickly, clearly still intoxicated by a cocktail of adrenaline and pride. He tells me about the incident in our interview a week later; how he’d gotten into a fight with another boy, Kyrone, because Harrison had ‘amped him up’. We are in the school library, drawing sharks at a table, but now Harrison stands up, physically retelling the altercation.

‘I kicked him in the head, I punched him. Oh no, I punched him there, I went dah!’ He swings at an imaginary Kyrone. ‘And then he got angry, kicked me there, kicked me in the muscle… like this bit?’ He lifts his leg and points.

‘In the calf?’ I clarify.

He nods. ‘In the calf, and then um, punched me there, like there,’ he points to his jaw and chest. ‘And a knuckle went there, and then I got pissed off and kicked him in the head. I kicked him in the head twice.’

I asked if he got in trouble when this happened, but Harrison wasn’t finished.
‘I… ah he like, when I kicked him in the head? I went like that, and then I fell on this side and then scraped my leg, down here?’

I said I remembered.

‘And then I had to go to the sick bay. And Miss, the one, the old teacher at room eight? Now she’s at room 15? yeah she’s like-’ he pants, hyperventilating. ‘And then she’s like wow your heart is beating like a cheetah!’ He laughs. ‘Yeah and I’m like-’ he gasps for breath. ‘I was angry and that.’

In describing this incident, Harrison positions himself not as a victim of bullying, but as the instigator of the fight, who ‘amps up’ Kyrone, initiates the first kick, and deals the final blows. He catalogues each moment of the fight, his injuries, and the experience of his bodily responses; the racing heartbeat and ragged breathing. He was an active participant in the fight and is active in its recall. This was an incident where he contested his vulnerability and won. Through his aggressive tactics, Harrison rejects victimhood; he is decidedly invulnerable, he is resilient.

Harrison is not the only boy to employ these aggressive practices of resilience, but they come at a cost. As well as the risk of physical injury, Harrison was regularly in trouble with his old school, and his behaviour may have improved his standing with his peers, but was incompatible with academic success. While Harrison had thus far evaded sanction for his behaviour at Tūrama School, other boys found themselves habitually in trouble, and I would regularly see the deputy principal walking across the school trailed by those who had been sent to him that day. Several boys were suspended during my time at Tūrama School for up to three days. As an accommodation for resilience, aggression could be a successful and culturally accepted strategy within the peer ecology, but it caused difficulties when conflicting with adult standards of behaviour and academic discipline.

Self-harm: a symptom or a practice of resilience?

Aggressive behaviour such as Harrison’s is one variable that would typically define ‘negative outcomes’ in a dimension of ‘prosocial’ behaviour often used as a measure of resilience in traditional research. This kind of research has been criticised for the implication that individuals who do not exhibit the ‘positive’ behaviour are not resilient, as measured by an ethnocentric social standard. In fact, a closer perspective, such as in Wang and Ho’s example
of the young women in urban Beijing, and the boys of Tūrama School, might reveal
behaviour such as aggression to be functional in specific domains, and therefore indicative of
an individual who has adapted to their circumstances in a way that works for them. Likewise,
it has been suggested that while dropping out of school is often seen as a ‘negative outcome’,
for some marginalised young people, this can be an adaptive strategy to sustain social
wellbeing (Dei 1997). Yehuda and Flory (2007) also argue that PTSD symptoms, the absence
of which have long been used as a marker of resilience, can be experienced by people who
nonetheless show great resilience in their capacity to cope with symptoms. PTSD symptoms
might actually indicate resilience when successfully managed, rather than a sign of less
resilient functioning. All of these examples point to issues with definitions of resilience that
rely on normative assumptions about pathology and adaptation.

In the same way that PTSD symptoms are constructed as indicators of pathology in
biomedical models of distress, self-harm is another phenomenon that would be considered a
‘symptom’ of ‘non-resilient functioning’ in much resilience research. Instead, I argue that
self-harm could be seen as another example of an embodied practice of resilience, at least in
this context. ‘Self-harm’ is the most recent in a revolving door of terminology⁶ used to
describe deliberate, self-inflicted injury such as cutting, burning, breaking bones, poisoning,
branding, scratching and picking at skin or pulling out hair. The term, which often eludes
clinical definition, excludes ‘socially sanctioned’ injury such as piercing or tattoos, or
harmful behaviour such as smoking or excessive alcohol consumption, and precludes suicidal
intent. Traditionally the purview of clinical fields, self-harm is still not well understood from
psychological perspectives which have tended to characterise the behaviour as histrionic or
stemming from poor impulse control (Steggals 2015). The inclusion of nonsuicidal self-
injury disorder (NSSID) in the latest edition of the Diagnostic and Statistical Manual of
Mental Disorders (DSM-V) (American Psychiatric Association, 2013)—not yet as a formal
diagnosis, but under the conditions for further study with proposed diagnostic criteria⁷—

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⁶ Also known over the years as self-injury, self-mutilation, deliberate self-harm, non-suicidal self-injury (NSSI) and parasuicide.
⁷ The addition, still controversial, was mainly to distinguish the behaviour from attempted suicide and from borderline
personality disorder, the only other disorder of which NSSI was a criterion (Wilkinson 2013). The diagnostic criteria include
engaging in the self-injury on five or more days in the past year, without suicidal intent, and motivated by seeking relief
from a negative state, resolving an interpersonal difficulty, or inducing a positive state, excluding ‘socially sanctioned’
behaviours (so as not to catch so-called normative practices such as tattooing, body piercing, or drinking alcohol in the
wrong net). Given that many of the criteria are based on individual motivations, it is difficult to assess how many of the girls
I worked with would have met the diagnosis, but most of these ten and eleven-year-old girls likely did.
nonetheless represents the next step in attempts to medicalise the phenomenon within a clean, clinical box.

Recently, sociologists have turned their attention to self-harm, unpacking behaviours like cutting not in terms of a natural and timeless category of illness but as an embodied, socially constructed phenomenon that is partially constituted by the psychological discourses that pathologise it (Adler and Adler 2011; Steggals 2015). While evidence of self-injurious behaviour is found throughout history, Steggals argues that the last two decades has seen the rise of a particular and novel form of self-harm, a heterogeneous psychopathology that has been produced out of punk/emo era bodily expressions of transgression and disenfranchisement and a medical framing which reinforces self-harm as a socially recognised idiom of distress. In other words, self-harm is a mental health issue because society historically said it was a mental health issue, and it has now become the quintessential expression of mental suffering, a ‘signifying wound’ that takes a ‘distinct and recognisable form in our contemporary cultural language’ (2015:1–3).

The prevalence of self-harm among children under 12 is not known in New Zealand, though the behaviour has been increasing among adolescents. Recent years have seen what has been described as an ‘epidemic’ of self-harm in Aotearoa, with 29.1% of girls, and 17.9% of boys in the most recent National Youth Health Survey of secondary school students (approx. age 12–18) reporting that they had deliberately hurt themselves in the last 12 months, up from 26% and 15.5% respectively in 2007 (Fleming et al. 2014). This is a particularly high rate when compared to other countries (Muehlenkamp et al. 2012), but mirrors the gender difference found in other studies, which early feminist analyses suggest reflect the result of gendered socialisation to externalise or internalise anger (Adler and Adler 2011). At Tūrama School, this was also a gendered practice, though I am not fully aware of its extent. I came to know of groups of girls in at least three classrooms, but in general this was a secretive practice that was known to other children but kept hidden from adults. Like some of the teachers, I caught glimpses when walking the line between the adult and child’s worlds, but it was a very partial view.

When Skylar comes to hug me and won’t let go, telling me she’s sad about other children teasing her about liking a boy, we wander into a conversation about shaving body hair and she tells me she cut herself with a razor, and then shows me her scars. Soraya was more direct, saying ‘look Julie’ before showing me the cuts on her arm. When playing paddle ball with Skylar’s friend Kaiyah, I catch sight of rows of parallel scratches and bruising on her
arm and when I ask about it she tells me she cuts herself. She runs off, but returns a little later, pointing to the bruises and asking me ‘what is this?’

‘Bruises?’ I ask.

‘No,’ she replies.

‘Well how did you get them?’ I ask.

‘From sucking like this’ she says, and demonstrates, twisting her arm to her mouth.

‘Oh, hickeys?’ I say.

‘Oh yeah, that’s it,’ and she runs off again.

The next day I go to their classroom after lunch to check if their teacher is aware of what is going on. The children are lined up outside their classroom, and seeing me, they call out and swing on the rail, leaning over to hug me or talk. ‘Look at this, Julie’ says Skylar, and shows me a long line of bruises on her arm. Kaiyah joins us and shows me hers. I rub Kaiyah’s arm and ask if anyone else knows about it. Kaiyah whips her arm away. ‘No, don’t tell anyone!’

She is whining, begging me not to tell. ‘I have to’, I say.

‘No! CYFs will take me away. They’ll think I’m being abused!’

‘They won’t think you’re being abused.’ I say gently. ‘They’ll think you’re unhappy at home.’

‘I am unhappy at home,’ she says matter-of-factly. ‘But they’ll take me away from my brother. They’ll take me away from my brother and I’m the only person he’s got. I don’t want to go without my brother.’

Self-harm is often described as a private or covert practice. But in this new iteration it is also, even when done in secret, a social practice, where feelings are experienced beyond the capacity of language to describe them and translated into a culturally recognisable, embodied expression of suffering. Kaiyah came to me seeking a label for her self-inflicted bruises, presumably not for her own personal knowledge, but so that they could be talked about and understood as distinct from accidental bruising. Soraya shared her wounds with me directly, telling me something she couldn’t express through language. In a world where adults may minimise or normalise children’s experiences of bullying and powerlessness, and conceptualise children as resilient and adaptable, it may be difficult for children to be heard in any other way.
When questioned by teachers, the girls said they were upset over a boy, and this became the accepted reason behind what tended to be seen as a ‘fad’ or attention seeking. While it may be true that some were upset about a boy, this does not explain why groups of girls across at least three different classes took up cutting at around the same time. Knowing the stories of the girls involved, including experiences of bullying, intervention by CYF, and living with hostile, alcoholic or abusive caregivers, not to mention other vulnerable experiences not recognised or validated by adults, it seems more likely that the story about a boy was simply an easy answer to questioning authorities. And while some of the girls may have been ‘copycatting’, as one teacher suggested, Adler and Adler (2011) point out that the documented ‘social contagion’ of self-injury represents the transformation of self-harm into a recognised, demedicalised social practice symbolising shared experience and social cohesion.

In its appearance as a sociological phenomenon, this form of self-harm does not easily fit with psychiatric models of a disorder of the troubled individual. Indeed Adler and Adler (2011) found a growing Internet community who espoused perspectives ranging from acceptance and normalising to embracing self-harm as a lifestyle choice or legitimate coping strategy that enabled functioning and represented a positive alternative to suicide. Wilkinson and Goodyer (2011) also point out problems with one of the DSM-V criteria which requires the NSSI to be causing clinically significant distress or interference across different domains of functioning, noting that this excludes large numbers of adolescents reporting NSSI because they tend to report their self-harm as helpful rather than distressing. This makes sense given that self-harm may accompany suicidal thoughts but is qualitatively different in that most people who self-harm do not wish to die, but use self-harm as a tool to alleviate anxiety, stress, or low moods.

Instead, self-harm may be viewed as practice of resilience—one which may not fit with adult ideas about what children should be doing, but which is nonetheless functional in the way it makes invisible suffering visible. A practice that is socially-sanctioned within the subcultures that foster it, self-harm enables that suffering to be expressed, shared, and validated. Viewing self-harm as a practice of resilience also illustrates the how resilience is built upon accommodations; self-harm improves functioning in some domains or in the short term, but at the cost of new vulnerabilities; the physical risks of injury and infection or the social impacts of scarring, stigma, and unwanted or negative attention. In this case, the school intervened and, unsure about what approach to take, initiated a chain of responses: involving the school social worker, contacting the families, and in the words of one teacher, explaining
the behaviour as ‘very naughty little girls’ who were only wanting attention and playing copy-cats.8 From a social ecology which created certain experiences of vulnerability, children were able to generate their own practices for coping, but a practice that was quickly closed off by adults, with little recognition of the conditions that had produced this practice or the potential need for alternatives to fulfill the same function.

Conclusion

This chapter has analysed children’s vulnerability and resilience within a socio-ecological framework that acknowledges both children’s agency and competency, and the environments that produce their vulnerability and constrain their opportunities. This ethnographic perspective suggests a shift in the lens through which socio-ecological approaches view resilience to include specific dimensions of resilience practices; coproduction, interconnectedness, and accommodations, and with the body as an important locus through which these processes can be traced.

Using social networks to illustrate, I emphasise that children do not enter an environment that pre-exists them, but actively coproduce that environment. As children like Ruby and Harrison experience the instability of relationships or direct threat of bullying, they exercise their agency in the form of strategic practices—the control of or aggression towards peers. Crucially, these practices shape the social environment, opening up new resources or creating new vulnerabilities that are experienced and negotiated by other children in turn. This interconnectedness of vulnerabilities and practices of resilience, seen also in the mutually constituting vulnerability of Mila and her mother, is a dimension of resilience that may be difficult to account for outside of ethnographic enquiry.

All of the children whose stories I tell in this chapter demonstrate resilience in different ways and in step with shifting circumstances they have little control over, but all of their practices of resilience vary in their success and incur costs; they are accommodations to their circumstances. Cassidee, who avoids fostering connections with peers she knows she may very well be soon leaving, misses out on the protective benefits that close relationships can bring. The tight control Ruby holds over her friends places children outside of the group into

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8 As this was at the very end of the school year, I was not aware of whether the effect of this disciplining was to halt the practice entirely or simply drive it into secrecy. (I, of course, would also no longer be allowed in on the secret, since I had made it clear that I was obligated to let someone know).
more vulnerable positions while offering resources to those within. Mila devotes great energy to the vigilant attention to her body and to her mother’s care. Other behaviours here that I have recognised as practices of resilience belong to that category of ‘atypical’ resilience (Ungar 2007) that would be instead labelled as ‘risks’, ‘symptoms’ or ‘negative outcomes’ by traditional resilience research. These too incur costs, but while Harrison’s aggressive behaviour compromises his ability to succeed academically at school, and puts him at risk of injury, it also allows a sense of agency and autonomy and self-efficacy he may not otherwise be able to access in this context. The self-harm practiced by many girls at Tūrama School cost them in terms of the disciplinary action taken by the school and the risk of unintended injury, but also gave them a sense of solidarity and social cohesion through partaking in this collective bodily expression that was likely protective, at least temporarily. The nature of resilience practices appears to vary by gender as strategies such as aggression have gendered implications—in this case, while it is more acceptable among peers for boys express aggression towards others, girls are more likely to turn violence towards themselves.

This is particularly relevant when considering how adults might resource children to support their resilience. Resources provisioned for the purpose of protecting children might be co-opted into strategies of resilience by some, contributing to a shift in the social ecology, which then opens up or constrains new resources or vulnerabilities. The clinic intervention, for example, appears to function as a catalyst for children’s coproduction of a new, collective vulnerability, one which plays out for some in hyper-vigilant self-care practices, and for others in denial or avoidance. Though this is a small ethnographic slice of life, the dimensions identified here—coproduction, interconnectedness, accommodations, and embodiment—could point to useful future pathways for the large-scale, mixed-methods studies currently leading resilience research.
Sitting around at our desks three weeks into the school year, I work quietly correcting spelling while listening to the chatter of the mixed-gender group of children around me. In general I found children’s conversation hard to capture; frenetic and stuttering, multiple threads tangling across each other, and individuals, catching words from the breeze, abandon one line of conversation to chase another, mouths fighting to describe a connection only their mind has made. As such, the chatter of children can sound like an incoherent layering of disparate ideas, interruptings and mumblings, a hubbub that is disregarded until it intrudes on adult conversation. Christensen describes how she had to develop an ability to ‘listen attentively’ to children and not be distracted by the interruptions of others when in a group (1999:75). Likewise, I found that I easily ‘tuned out’ children’s talk in such situations, since it required a special effort to listen and often lost me along the way.

On this day, the conversation clarified for a few moments around a particular topic that captured everyone’s attention: what are you scared of? The answers fought to impress: aliens, crocodiles, snakes. I noticed with interest that the things named were not things that the children had any real experience with. ‘I’m not scared of anything’ came one boast.

‘You’d be scared if your mum died,’ came the retort. There was general agreement; a parent dying was a scary thought.

Childhoods in Aotearoa are not homogenous and the degree to which death appears in childhood, both in fantasy and in reality, will likely vary alongside children’s conceptions of death. For children at Tūrama School encounters with death are common. For many who are Māori, wide whānau connections mean that journeys to the marae for tangi, or funerals, are frequent, often extending three or four days at a time, and for many children, experienced like a holiday (Jacob 2011). For the Pasifika children, too, the passing of relatives may mean a trip to the islands, to Tonga, where funerals are a preeminent social occasion and key locus for enculturation of the young (Kaeppler 1978), or to Samoa, where the dead stay alongside the living in front yard graves (Havea 2013). In Auckland, too, family homes may be converted into ‘little marae’ for Māori (Gagné 2013) and Pasifika lives celebrated with ritual even more traditional than in the home islands (Havea 2013).
Many children at Tūrama School also experienced the death of a close relative, including five children who lost a parent during my study period. Often, these children would quietly disappear from school as they were rehomed with whānau in another part of the city, leaving behind classmates who named what had happened with wide eyes and reverent voices. These children are socially marked, spoken of with hushed voices and respectful distance. They have experienced the far edge of child vulnerability, that place spoken of in fear across the classroom desks. That vulnerability is found in not just the daunting grief of the loss of parent, but the social upheaval that follows for a child who must start a new life not of one’s choosing.

Child death frequently appears in ethnography, where childhood mortality often forms the backdrop to ethnography of poverty stricken or war torn communities—the ‘unnaturalness’ of child death used to emphasise the direness of conditions through statistics or vignettes. Nancy Scheper-Hughes’ (1993) ethnography of shantytown Brazil paints a community so saturated with child death as to cleave the most primal bonds of mother and infant. Yet children’s experiences of the death of their siblings, or their own close brushes with mortality, remain outside of the picture.

Only a small library of anthropological works represent the perspectives of children themselves on death. Significant works by Myra Bluebond-Langner (1978, 2000) have focussed on chronically or terminally ill children in a North American context, demonstrating how children acquire knowledge of their prognosis or come to understand that their sibling will die. Based on research in Dehli, Veena Das (1989, 2015) reflects on the experiences and expressions of children who have witnessed violent deaths or coped with the loss of a parent. Ross Parsons (2012) combines psychotherapeutic and anthropological understandings to write of the ambivalent liminality of HIV positive children enduring poverty in ‘the waiting room of death’ in Zimbabwe. From different corners of the world, these scholars trace the way that death shapes the social world that children live in, just as it is shaped by adult notions of how children’s encounters with death should be managed. Yet for the most part, death in this literature is represented as an anomaly, even when common, undermining normal, carefree childhoods, and leaving children to embroider its extraordinary tragedy in repetitive symbolic or embodied expressions. With the exception of Das’ (2015) portrait of a child renegotiating changes to the family structure as a result of illness and death, these are portrayed as transgressive deaths, not a death that touches childhood in the everyday.
The limited explorations of death in everyday childhood perhaps reflects what scholars have claimed to be a denial or general suppression of death in contemporary Western culture (Becker 1973; Ariès 1981), where despite near saturation of death imagery in the media, personal experiences of death are kept hidden or ‘invisible’ and marked as taboo. This death, James Green (2012) adds, is seen as particularly inappropriate for children, whose lives are meant to be innocent and happy, free from adult burdens. Once commonplace, over the last century death has come to be seen as out of place in children’s everyday in the recent West (Kastenbaum and Fox 2008), and viewed as a threat to the social order (Bluebond-Langner 1978). This is well illustrated by the ‘mutual-pretense’ of Bluebond-Langner’s (1978) dying children and their parents: planning Christmases that would never arrive, continuing with school, and maintaining the socialisation of the child for an adulthood that all knew would not come. Discussions of death and childhood therefore often beg in with the Western premise that death ought not to be part of children’s lives, analysing the way adults and children alike make sense of this anomaly.

However, death is part of children’s everyday lifeworlds, though their personal experiences of death can be limited. From children’s films, where death is seen 2.5 times as often as adult films (Colman et al. 2014), to religious stories, to violent toys and video games, to classroom discussions (at Tūrama School, a full term on World War One), death is already central to children’s cultural systems, structuring everyday meaning and practices and enacted through play. Pioneering studies by Nagy (1948) and Anthony (1972) in early psychology showed that death-related phenomena often appear in children’s thinking within their everyday lives, and children had their own questions and theories. In conversations with Suffolk young people, sociologist Sarah Coombs (2014) demonstrates the extent to which young people absorb and incorporate these cultural scripts into their constructions of death: the romantic death, the heroic death, the violent death. From a childhood studies research project on children’s personal lives, Hayley Davies (2017) found that children often referenced death in the telling of their biographical narratives, whether this be the death of a grandparent or pet, or the anticipated death of parents, and memorialised bereavements in sensory and material ways. For Māori and Pasifika children, the presence of death in the everyday is even more overt, and their personal experiences of death more frequent than their Pākehā contemporaries. What role then, does death play, and what meanings do children construct from death in the everyday?
Small Talk of Death

I did not set out to learn about children’s perspectives on death. I did not raise death as a question in the interviews I held after school with children, parents, or teachers, and I did not explore children’s experiences of tangi. However, I found that my field notes, interviews, and even children’s drawings were saturated with references to death—unsurprising, given the proximity of death in these children’s lives, and revealing of the salience and relevance of death to childhood. Children catalogued, memorialised and constructed relationships with the dead, concocted understandings of the afterlife and experimented with mortality through risk. I cannot offer here a child ontology of death, or comment on their understandings and knowledges of death—a topic that has been considerably discussed within psychology and counselling fields (e.g. Anthony 1972; Cuddy-Casey and Orvaschel 1997; Nagy 1948). Instead, this is an analysis of the way that death is woven into the discourses and practices of everyday life outside of death events and rituals, the appearance of death in life’s little conversations, and what this can tell us about childhoods lived in the classroom; at the kitchen table; on the playground. This differs from studies which set out to ask about death, and instead stumbles across the ways that death creeps unsummoned into everyday childhoods.

In children’s ‘small talk of death’—how children express death, and how children use death to express other things—can be found the social meanings children construct around death, but also the experience of childhoods as shaped by broader political, economic and social forces. As such, I argue that the meanings of death for children are shaped by their social position as children, a possibly universal process of apprehending this intriguing idea that existences will end, coupled with a structural vulnerability which inoculates the interpretations of those unknowable deaths with threat of social displacement. However, concepts of death are also coloured by the particularities of children’s social circumstances, and so for the children at Tūrama School, their death talk also reflects plural cultural frames and a structural violence that mediates their experience of death. Children’s understandings of death are shaped by their experience of life, and so children’s death talk can also function as a window for understanding childhood.

As co-constructors of their own cultures, but also embedded within adult worlds, the way children appropriate and reproduce their own versions of death can be usefully conceptualised through William Corsaro’s (1992) model of interpretive reproduction, where
children spin their own webs of meaning over the scaffolding of adult institutions. As described in chapter one, Corsaro breaks away from theories of socialisation which position the child as individually and privately acquiring the skills, knowledge, and culture of adults, instead viewing childhood socialisation as a collective, social process, occurring within peer cultures as much as with adults, and reproductive rather than linear. My analysis, therefore, begins in children’s peer talk, where snippets of experience are collectively patch-worked.

**Death in Cultural and Structural Context**

Eleven-year-old Arya was eager to be interviewed and as soon as I turn on the recorder she begins talking and doesn’t stop, one thought tumbling into another, punctuated with bright-eyed laughter and the rising inflection common of Kiwi cadence. She begins by telling me about her family’s history of gang violence, but her stream of consciousness leads into a cataloguing of deaths in her family, a theme she keeps returning to annotate again and again.

‘And then my cousin, he wanted to give the Crips gang a hiding, but they told him that they should just leave us alone? We had like, bad people in our family? That got rushed- my cousin got… beaten last week? And my papa died the year before, like 2012? And our papa died? Well my other papa died on my dad’s side, ages ago, I think it was 2010? And it was heartbreaking because he was a good cooker and- and- we loved him and we gave him everything, like he wanted?’

Disoriented by the shift in topic, I ask, ‘did he die because of the gangs?’

‘Oh, no, he died because he had a back injury. Oh, sore back.’

‘He died from a sore back?’

‘And he was sick,’ she adds, and continues, ‘that's the same as my little sister. She’s like, eight now, and she passed away in 2007. And my cousin, oh, we had a big-as tangi and funeral? Because everyone liked her, and my auntie, that was her- that was her godmother? And that’s the one that- she loves me, now? But she still- she still loves her, like her daughter, like she wants her to be her daughter? And- and my family was so sad. And that was my little sister, she’s- she turned eight, in… April. On the 23rd.’

‘Do you guys do anything for her birthday?’

‘Yes, we went to go clean her grave and put flowers and stuff. And… we sing- sometimes sing a song to her. When we go past.’
Later in the interview Arya returns to her topic, adding further infant deaths that occurred before she was born, and the death of another papa in 2014. The way children like Arya talked about the deaths of their relatives echo some universal fascinations with the nature of existence and its cessation, the continuation of relationships after death, and some specific features of childhood. However, children’s death talk is also shaded with the particularities of their circumstances which add the nuance of experience. For children at Tūrama School, these particularities include a structural violence that burdens whānau with earlier and more frequent death. In addition, these are children of plural and syncretic cultures, and so incorporate multiple frames of death into their own understandings. Children experience, for example, the way that Western medicalised frames of death shape everyday life into a series of risk preventions (McIntosh 2001), from road crossing patrols at school, to legislation mandating ‘child resistant closures’ on medications and chemicals. Likewise, Westernised medical institutions invoke death in the management of illness, which Tūrama School children absorb through health promotion warnings about the deadly threat of rheumatic fever. Meanwhile, predominantly of Māori and Pasifika descent, many of these children have grown up in cultures where death is not out of place in children’s lives, dying not cordoned off into hospitals nor old age. Instead, death is axial to life, the boundaries between living and dead blurred, and children themselves may be considered intermediaries to the spirit world (Counts and Counts 1985; Gagné 2013; Metge 1967).

For example, care and connections with the dead are infused throughout te Ao Māori, or the Māori world, and vice versa, as expressed by Māori writer Harry Dansey: ‘we cannot think of the dead without reference to the living’ (1975:174). The depth of relationship forged from shared, even ancient ancestry, is tied to a deep emotional expression for even those little known to the mourner (McIntosh 2001; Rosenblatt 2011; Sinclair 1990). Death is also embedded in the architecture of Māori social life, where marae, or traditional meeting places are explicitly designed for the hosting of many visitors for tangihanga (Gagné 2013; Salmond 1976). This centrality of tangi to Māori life is far from the ‘invisible death’ that Philippe Ariès (1981) saw as marking contemporary Western cultures. Very little has been written about children’s perspectives of tangi, although Jacob’s (2011) research with Māori parents includes the recollections of adult participants who compare childhood experiences of attending tangi to going on holiday. For these Māori, tangi were not unusual, but remembered as a break from the normal everyday grind, a chance to see relatives and play with cousins for days on end, free of adult responsibility and relatively unsupervised. Depending on the
closeness of the deceased, the business of grieving could be secondary to the excitement of the trip and reunion with whānau. This was also a familiar experience for the Māori children of Tūrama School and talk of attending tangi formed a part of everyday conversation.

Yet while culture makes meaning of the dead, encounters with death are also mediated by the structural. Living at the extreme end of socio-economic disadvantage, Tūrama School children’s experience of death is again amplified. To understand how death articulates with context here, I use the related concepts of structural violence and structural vulnerability. Commonly attributed to Johan Galtung (1969), structural violence is a concept used by anthropologists such as Paul Farmer (2003, 2004) and Nancy Scheper-Hughes (2004, 1996) to bring a critical perspective to health inequalities. Structural violence refers to the way that sets of historical and economic conditions, when protected as the status quo, result in physical or spiritual harm, disease, oppression, and premature death (Farmer 2003). While the effects of poverty, racism or gender inequalities can be misrecognised as misfortune, structural violence conceptualises systems—and the people who uphold them—as perpetrators of harm. Quesada and colleagues (2011) propose ‘structural vulnerability’ as an extension of structural violence which explicitly includes cultural and idiosyncratic sources of structurally mediated distress, such as symbolic taxonomies of worthiness or discourses of normativity. Children, because of their perceived vulnerability, incompetence and low status, are also rendered structurally vulnerable through society’s adult-centric organisation, often encoded in legislation, which separates them from the adult world, constraining their political power and limiting their voice (Frankenberg, Robinson, and Delahooke 2000).

The appearance of death in the cultural milieu of Tūrama School can be seen as both a reflection of a structural violence that mediates the degree of children’s experience with death, and a structural vulnerability that produces a particular experience of childhood. As a result of economic inequities, institutional racism and colonisation, Māori death rates tend to be higher than those for non-Māori at all ages, though the gap is narrowing, and life expectancy at birth for Māori is about seven years lower than for non-Māori in New Zealand (Statistics New Zealand 2015). Pasifika peoples in New Zealand have life expectancy about 1.5 years higher than for Māori, but are still also well below the rest of the population. Infant mortality rates are higher for Māori (0.65%) and Pasifika (0.62%) than for the general population (0.45%) (Statistics New Zealand 2015), and 2013 data show Māori suicide rates were 1.6 times higher than the overall population, and Māori youth suicide rate was 3.1 times the rate for non-Māori (Ministry of Health 2016).
The experiences of the children living in the community surrounding Tūrama School reflect these statistics and the death of a close relative was not uncommon. Aside from the passing of many grandparents during the study period, five children in my cohort (two of them brothers) lost parents to illness-related deaths. The deaths of each of these parents affected not only their children, but also their nephews, nieces and cousins attending the school. Some children disclosed personal knowledge of a relative who had died by suicide, while two ten-year-old boys described to me the phenomenon of suicide ideation in detail, suggesting great familiarity (‘like having a bad life and everything has been… bad in their life, they could say I want to kill myself. Then change their mind and think about their family, and what that will do. And then they say oh nah’). For eleven-year-old Pikau, the ideation was a personal experience, as she disclosed to me how she had thought of killing herself after years of being bullied.

As well as the structural position of Māori and Pasifika which brings death earlier, the structural vulnerability of children in society gives these deaths special meaning. As in many Western countries, children in Aotearoa are one of the most legally and socially restricted groups in society, with limited mobility, political power, or earning ability. Although children’s smaller body size leaves them more vulnerable than adults, their dependency in these contexts is to a large degree created by adults who are concerned with their care and protection. In other societies, even very young children can move independently through their community, access material resources, and care for each other in peer groups, rather than rely exclusively on adults. In some circumstances street children, for example, can obtain better nutrition than children who live at home (Gross, Landfried, and Herman 1996), indicating that dependency on adult care is to a certain extent structural, rather than only biological. Children’s near complete dependence on adults in Aotearoa therefore, is less about an innate vulnerability, and more the product of their structural vulnerability in an adult-centric society which often marginalises children.

Children’s structural vulnerability has two implications relevant to this discussion of death. Firstly, in rendering children so dependent on adults, the threat of death, particularly of a caregiver, represents a threat to the social life of the child. When parents fall ill or die, dependent children must be placed elsewhere in the whānau, a dislocation of relationships, home and often school that can be a disorientating addendum to grief, even though children’s belonging is more fluid in Pacific kinship than for Pākehā (Metge 1967; Morton 1996; Ritchie and Ritchie 1979).
Secondly, the adult-centric distribution of power in society often means that children are invisibilised in culture and policy. Subsequently, children may be exploited for adult agendas, with little regard for what this means for children. This can particularly be seen in health promotion messaging which links children and medicalised, Pākehā frames of death in order to motivate parents, but inadvertently influences children’s conceptions of death and subsequent practices, as in the rheumatic fever campaign. The experience of structural violence and structural vulnerability for the children of Tūrama School can both be observed through analyses of both how children express death, and how they use death to express in everyday life.

Expressing Death

At morning tea one day, I sit with Ruby and Soraya, our backs against the classroom wall at the edge of the playground. As we watch a group of girls and boys kicking a soccer ball around on the field, I tune in to their talk, which has taken on a sombre tone. ‘My nephew died,’ says Soraya, gap toothed and with a smile that curls up at the edges. ‘My little four-year-old nephew, you know? Because he wasn’t eating anything.’

‘My cousin died,’ says Ruby, her eyes big.

‘Everything he ate up, he threw up,’ Soraya continues. ‘Vegetables… meat…’

‘-My cousin was 16,’ adds Ruby.

‘…lollies… everything, he threw up.’ A stray ball flees towards us, and Soraya intercepts it with the edge of her foot, and boots it back towards the field.

‘Did you know he was going to die?’ Ruby asks.

‘No,’ Soraya answers. ‘We didn’t know he was sick until we took him to hospital. And then they sent him home, but then he died.’

‘My cousin hung herself,’ says Ruby. That’s really sad, I respond.

‘Oh my god!’ Soraya interjects. ‘That’s the same- not the same, but that’s the same thing as my little nephew. But he died in hospital. ‘Cause he was getting fed the right things but he couldn’t breathe when he ate it. Like every time he ate something, he would just stop breathing.’
This ‘small talk of death’ is woven through children’s desktop chatter, in the sharing of global and personal news on the classroom mat, and in playground banter. These small conversations are where children engage in a process of collective bricolage, pooling their experiences of death to make sense of its place in their world. Ruby and Soraya ‘name’ the whānau members who have died, an intensive cataloguing of deaths which is usually accompanied by the manner of death, if known. Children often included dates in their catalogues, and the age the sibling would be; Arya refers to her sister as ‘eight now’. This is a sharing and comparing of experiences in order to apprehend death, but children also register who knows what by indexing who was affected (‘Liam’s mother died. And she was Trystan’s auntie’). While I contribute a comment on the sadness of the death, as teachers tend to do, Ruby and Soraya do not talk about what feelings or meanings they associate with the death. Instead, they add to their catalogue, a practice which takes on a tone of one-upmanship, using experience of death to compete and impress, and drawing comparisons; Ruby’s cousin’s death by hanging was ‘the same’ as Soraya’s nephew who couldn’t eat. Claiming a close experience with death could thus function as a form of social capital, impressing others with the reminder of their vulnerability and the knowledge of that which all children will experience at some point, but which some have not yet faced. In an interview with two nine-year-old boys, I hear the following snippet of conversation:

Whetu: None of my family has ever died yet. Only my nan. My mum’s mum.

Jackson: My nan died two years ago.

Whetu: My nan died four years ago.

Rehearsing cultural scripts for talking about death through sombre tones, in this talk children situate themselves in wider networks of relationships which continue after death. At the same time, the comparisons and repeated indexing of death also marks these deaths as a salient part of children’s experience.

Children’s expressions of death therefore express their experience of life, one where the deaths of family members echo with threats about those who could die. ‘Whose mums and dads died in this group?’ one boy asked a group of classmates during a brainstorm of what it means to be a child. ‘If you’re young, and your mum and dad die, who would you live with?’ ‘Your grandparents,’ another child replies.

‘Yeah your grandparents.’
‘Or your brother and sister if you have them. If they’re older.’

What it means to be a child, then, is to be dependent on networks of care for social survival.

Death in the classroom

On the first day back from the spring holidays, after the usual Monday assembly, Mrs Steven’s class settles on to the mat to start the day. As is their routine, Mrs Stevens asks, ‘what news do we have today?’ The children scramble to name the local and global headlines they’d picked up over the holidays. New Zealand has made the quarter-finals of the Rugby World Cup, and evidence has been found of water on Mars. ‘Lots of murders,’ one child calls out.

‘Yeah that boy got murdered!’

‘Yes I think his funeral is today,’ Mrs Stevens acknowledges. The disappearance and subsequent discovery of a ten-year-old’s body in the South Island had captured the attention of the nation for the past week. ‘That boy was murdered, Julie,’ a girl said to me.

‘A lady died while she was feeding her baby and her baby died too!’ another child adds, and Mrs Stevens pulls open her laptop and finds the story from earlier that week. ‘How did she die, Miss?’ children ask, and Mrs Steven’s reads from the article. ‘She collapsed and died of bronchitis.’

‘What’s bronchitis?’ Mrs Steven’s explains about infections in the bronchial tubes; this is how much of the learning happens in this class, through informal dialogue on the mat.

‘My auntie died Miss! Aye miss! Aye, my auntie died!’ Eleven-year-old Trystan, who was lying on the ground, leaps to his feet.

‘Oh yes, she did,’ Mrs Stevens realises, meeting my eye.

‘She died of new- pneuma- pneumonia,’ Trystan wraps his tongue around the word. ‘She didn’t go to the doctor.’

‘And she was Liam’s mum!’ Another child adds.

‘What?’ I hadn’t noticed Liam’s absence until that moment. Quiet and small, Liam had joined the class only a few months ago.
‘Yeah, Liam’s moved to Ferndale,’ Mrs Stevens tells the class. Turning to me, she adds, ‘it’s a shame he’s not here so we could support him with his mum dying. He didn’t want to leave Tūrama.’

Figure 9. ‘My auntie died, Miss!’

Death, then, makes a regular presence in the classroom, not only for these children who are directly affected by loss, but also for those around them who bear witness to and imagine as their own the experiences of their classmates and the boy in the newspaper. After rugby news, children tended to share the death stories, a cataloguing of global deaths to mirror the real ones they encountered. But their structural vulnerability is brought into relief this day, where the death of Liam’s mother was followed by his abrupt disappearance from the class he liked and the teacher who cared for him.

Trystan was clearly impacted by the death of his aunt, which he brought up with me on several other occasions. However, the expressions of children who had lost a parent were quieter; Teuila, tells me that her dad is—present tense—an artist who taught her to draw, and sits silently as her teacher reads to the class newspaper articles about his premature death. In my interview with Cassidee, I invited her to draw for me a picture illustrating either
something she enjoyed doing at school, or a time when she found things tough. She would not let me see her drawing in the interview, frantically covering it with her arms, and asking that I only look at it after she left. When I remembered to pull out the drawing a few days later, I found a depiction of her father’s death (Figure 10).

![Drawing of a father's death](image)

Figure 10. ‘When My Dad Died’ by Cassidee, age 10. The pink writing is presumably referring to the experience of being interviewed.

Once death has struck so close, children’s expressions turn away from its apprehension, and towards commemorating, memorialising, and configuring relationships with the dead.

Expressions of death; death as expression

The way that the children of Tūrama School talk of death is consistent with how death is expressed by children in many other contexts in academic literature, where ethnographic vignettes show children engaging directly and expressively with death encounters. These expressions can often look quite different from those of adults in the same community; in the
context of Tongan childhoods, for example, Morton describes the aftermath of the suicide of a neighbour boy, where adults are distressed and quiet, while ‘children were excitedly chattering, telling me the details of how he had been found, what his face looked like in death, how it was originally thought he must have been electrocuted, and how his mother was in deep shock, rocking on her bed and calling out to him as if he were still alive’ (Morton 1996:241–42). When adults began to recover from the shock, they discussed the death only briefly to agree on the motivations behind the suicide, before moving on to planning the funeral and positive recollections of the boy and his mother.

Likewise, among families of children with cystic fibrosis, Bluebond-Langer (2000) found that although parents and children rarely talked to each other about the prognosis, children’s talk is littered with references to death, which increase as their sibling’s condition progresses. While parents avoided thinking about or discussing the prognosis even as their child’s condition deteriorated, children’s talk belies a preoccupation with dying. However, children’s expressions are not limited to the verbal. In her earlier work with terminal leukaemia patients, Bluebond-Langer describes a proliferation of death imagery emerging after children came to understand that they were dying; a child who used to draw birds and flowers began to draw crucifixion scenes, another made only ‘turtles for people’s graves’ (1978:185) and buried her paper dolls in tissue boxes, and cohorts of children asked to be read the chapter of Charlotte’s Web where Charlotte dies. Behaviourally too, children expressed their impending death, for example through a concern with the passage of time: admonishing parents not to ‘waste time’ and becoming angry when people took ‘too long’. Das (1989) also describes how Dehli children who witnessed anti-Sikh riots expressed the death of parents in embodied ways; one child mimes the hanging of his father, while others are haunted by nightmares of ghosts. Vivid and direct expressions of death are therefore found among children in many contexts where death has been experienced.

These vignettes are used to reveal children’s knowledge of their, or their sibling’s prognosis, how children make sense of the death and violence they have witnessed, and how they draw together fragments of adult conversations to weave their own coherent narratives in a collective reconstructing of events and experiences. In unpacking Tūrama School children’s references to death, it is apparent that children are not only expressing knowledge of death or fear or sadness, nor just engaging in a collective bricolage to make sense of this death concept. The children also spoke of death not in reference to death itself, but in order to invoke the connotations of death to express other experiences. Death was not only expressed,
but is used to express. So when Ruby and Soraya tell me that some people who are cyberbullied ‘get to like, dying’, they are telling me about the seriousness of cyberbullying, as well as the seriousness of death. When 11-year-old Navahn tells me about the gang his family are involved in, he tells me ‘they have heaps of fights, they die’ to convey the degree of violence. And when nine-year-old Hinemaia says strep throat ‘could stop you from breathing, and that could kill you too’, she uses death to explain the threat of a sore throat. Children’s talk of death, therefore, is important for what it says about life.

Expressing illness with death

Children’s appropriation of death terminology to communicate ideas about other things can be most clearly seen in their expression of illness. The bundling of death with illness is in part because death in this community is often caused by illness, though death could also be used to convey ideas about bullying and violence, as above. However, to speak about illness also requires a specific lexicon, one that children are still acquiring. By contrast, when parents speak of illness, the language employed is quite different from that of children. Anna, in telling me of her son Anton’s illness history, draws from an extensive and nuanced language of sickness, acquired through her years of parenting experience. When Anton was born he was ‘meconium’, ‘stressed because he was overdue’ and ‘he wasn’t breathing’. She describes the efforts of the doctors, the effects on his throat and lungs, how his temperature of 40 could lead to a seizure, and the work she did to steam his excess mucus. This is not a language that Anton shares. When he tells me about ‘one of my bad sickness’, a spider bite, he tells me that his mum took him to the doctors, and describes what the doctor said: ‘Your son got poisoned, but I don’t think he’s gonna like die or something, it’s just like, you know, he’s gonna be sick.’ He adds, ‘and my greatest fear was like, dying.’ After telling me about how the spider bite ‘restart my memories’ and ignited a new interest in soccer, Anton returns to the theme of death:

‘I was in hospital just for the great white tail spider, but, lucky… he didn’t said, you got a poisoning and you’re gonna die. Lucky he didn’t said that. But he says he got a cure, and that. And then, he- he like had a needle put in my like, I think like right here?’

‘Oh he put a needle in your side?’

‘Yeah. And then right before, he said, you’re not going to die.’
Lacking the vocabulary of symptoms and treatment, Anton emphasises his experience in terms of proximity to death—Anton feared he might die, but luckily the doctor said he wasn’t going to die, such was the seriousness of his sickness. When I ask him how come he was thinking about dying, he grapples for the words to express the experience of his body in that moment:

‘It kind of felt like, I kinda feel like my heart- my heart beating was like, like, slowing down… Wait, I was actually- I was actually ill, but- a little ill, so- it was still- it was like going slooooww… and then going faster and then going slow and then going faster. It’s kind of... It’s kind of scared. And like, and I was like um… was my heart going to explode or something.’

The language of death, therefore, can function for children as an effective shortcut to communicate the seriousness of an illness experience, a way that children can express and be understood by other children unfamiliar with the vocabulary of arrhythmia or anti-venom injections.

Similarly, death is the focus of the story of Te Rerenga’s birth, as narrated by her and her sister Nga-Atawhainga in the middle of a discussion about illness experiences at their kitchen table. After asking her mother whether she had been sick as a baby, nine-year-old Nga announces:

‘My sister nearly died when I was a baby.’

‘Yes no, she had a struggle to life, Rere did,’ her mother Tūmanako agrees.

‘She nearly died when she come out. Mum told us.’

‘How does that feel?’ I ask eight-year-old Rere, whose face is ringed with chocolate from the biscuits I brought.

‘Like, the same,’ Rere answers, I think meaning she feels the same as she always does, that how she feels is not the important thing. She continues. ‘My mum nearly died when she got-’

‘-Mum got a fever when I was having her,’ Tūmanako interjects, switching narrative perspective halfway through her sentence. While Rere attempts to continue, Tūmanako narrates a parallel story, this one told through symptoms and treatments and their effects.

‘Oh, and I nearly died-’

‘And then because we had an inexperienced person trying-’
‘-I was sick when I was four or five-’
‘-to give me an epidural, they nearly paralysed me-’
‘-when I was little you took me to the hospital-’

‘-because you were too big to come out I had to have a caesarean section. I tried to tell them that, ‘cause I’d been in labour for five hours, I was like look, last time it only took me an hour for the whole thing, I'm telling you I can’t do this, please listen, you’re going to kill me,’ Tūmanako finishes, laughing. ‘She didn’t have a good start to life, Rere.’

Interestingly, as her daughters, lacking the language of labour and caesareans, refer repeatedly to death to convey to me the gravity of the event, Tūmanako’s only reference to death is also deployed in another moment of misunderstanding. ‘You’re going to kill me,’ she laughs now, but at the time communicated what Tūmanako’s body knew, but couldn’t express in any other way to the ‘inexperienced’ doctor.

With less experience of verbal communication, particularly the vocabulary associated with exceptional illness events, children therefore draw on their own categories of concepts to signify the most serious, the most frightening of events. Along with death, Anton and Rere both make reference to the hospital, another signifier of a serious medical condition. For the children at Tūrama School, the hospital is an important symbol of sickness, differentiating notable sickness from the everyday. In one lunchtime conversation, children discussed hospital visits, asking me how many times I’d been to hospital and calculating the times they went to hospital and their injuries.

The use of death language in their everyday talk therefore also speaks to what death means for children. Adults, with longer experience, may have more of a sense of life’s rhythms than children, who are still calibrating their sense of death’s relative frequency. Adults can suffer greatly in bereavement and face social disruption, but with their greater agency and autonomy are less socially threatened by the death of relatives than children are by the loss of caregivers. For children, death means the deepest and most insecure grief, as they are tossed into the waves of a social upheaval, with little control over where and with whom they may land.
Children’s talk of death in this context reflects their own structural vulnerability, the structural violence that makes death a common occurrence in this community, and cultures that make meaning from those deaths. However, their conceptions of death are also shaped by Pākehā medicalisations of death, how, as McIntosh (2001) describes, under the rise of modern technologies and medicines, mortality is deconstructed into a series of discrete problems, manageable through ‘taking control’ of the body, adopting healthy practices and abstaining from the unhealthy. When death comes to be seen as solvable, then life becomes a series of risk preventions, as McIntosh puts it, ‘an attempt to write death small’ (2001:247) which paradoxically leaves death imprinted over life in seatbelts and alarms and expiry dates. Public health campaigns fixate on death as the enemy, even when, as is the case with rheumatic fever, death is the least likely outcome. The effect of this is to semiotically link illness to death, and health care, even for children, becomes constructed in terms of death management.

‘And what about cancer, what do you know about that?’ I ask nine-year-old Hinemaia, after she includes cancer in her list of illnesses. ‘That it could actually kill you,’ she replies; her nana died of cancer. While we are looking at the crosses that have been put out by the school playground to commemorate the soldiers who died in World War One, 11-year-old Pikau tells me about her auntie who died, an auntie who had been very sick because she had a colostomy bag. Ten-year-old Chloe tells me how her church played videos of children in Pakistan who are ‘really sick and if one of the boys there gets one more drink of dirty water he’ll die.’ Cancer is memorable because it is deadly, the fates of soldiers in war are linked to a colostomy bag, and children in Pakistan must be close to dying if they are sick. In this way, sickness and death are, for Tūrama School children, entangled as each constitutes the other. This coupling of death and illness likely comes in part from those parents and grandparents whose death was attributed to illness, like Trystan’s auntie who apparently died of pneumonia. A great weapon of structural violence is the way deaths become naturalised, ascribed to their proximate causes rather than the systemic inequalities that left those bodies vulnerable to illness in the first place. However, recent media campaigns in New Zealand which both involve children and draw on death as a motivator contribute to the meaning of death for children, as well as inadvertently target children with powerful, frightening messages that are meant for their parents. The One Heart Many Lives campaign, which aims
to ‘inspire Māori and Pacific Island men to get a heart check’ (PHARMAC 2017), includes an advertisement aired on Māori Television showing a young girl speaking to her father about how important he is to her and to the protection of Māori tikanga [customs] and taonga [treasures]. ‘But I’m too young to visit you in the cemetery, Dad,’ the girl says, sitting in tears outside the whare nui. ‘See a doctor, and get your heart checked. I love you dad.’ This advertisement uses the tears of a grieving child to motivate fathers to have their heart checked, but also likely impacts children who identify with the young girl and fear the loss of their fathers. Aired on the national Māori television channel, this campaign also speaks specifically to Māori children. Similarly, the rheumatic fever campaign, which is intended to target Māori and Pacific parents, in fact speaks directly to children. Many of the television and poster advertisements feature children, telling their stories with explicit links to death. ‘Last year, I almost died,’ a young boy tells the camera over a solemn piano motif and ambulance sirens. Death, as it is portrayed in these advertisements, is not the gentle, spiritual crossing over to te Ao Wairua [the spiritual world] that children hear narrated at tangihanga, but one of panicked lights and sirens, alienating surgical theatres and thick scars that witness the ripping open of a body, a close evasion of a terrible fate. This death is used to connot the most extreme feelings of fear or anxiety, and in doing so, constitutes a terrifying new version of death. Importantly, although this is a framing of illness in terms of death, these advertisements feature Māori and Tongan children, and as such, directly target not only children, but specifically Māori and Pasifika children.

If death represents a powerful tool of expression for children, then portrayals of child death in the media have a particular, unrecognised potency. While children’s low status and presumed passive role often renders them invisible in policy and planning, paradoxically, calls to ‘think about the children’ are often used for the purpose of mobilising adult action, without regard for children’s interpretation of this messaging itself. The invisibility of children in society means that the Health Promotion Agency did not consider how they would view these advertisements (personal communication). However, the effect of the rheumatic fever campaign on the children of Tūrama School is very evident. Children would recite the campaign messages verbatim; ten-year-old Te Kapua, placing himself in the shoes of the boy, narrates: ‘My brother almost died. It started with a sore throat.’ Half a world away, Jean Hunleth (2017) describes a similar response among Zambian children to orphanhood.

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1 The campaign was led by PHARMAC (New Zealand’s pharmaceutical management agency) from 2008-2013 and is now held within local communities.
2 Main building of a marae, where guests are accommodated and hui (meetings) held.
discourses propagated by adults to draw attention to HIV and related illnesses. Repeating slogans they heard on the radio and in school, even children who technically already were orphans expressed fear of becoming an orphan. Children constructed a meaning of orphanhood with similar valence to the meaning of death for Tūrama School children; connoting loss of social relationships, care and belonging.

In chapter four I introduced Marielle, who misunderstood her two bouts of strep throat to be ‘rheumatic fever’, which, for her, was characterised by the experience of breathlessness; a symptom the nurse later attributed to anxious hyperventilation. Marielle’s interpretation of her experience seems to have been directly shaped by the rheumatic fever television campaign, which she repeatedly references, along with her fear that she was going to die. After nearly crying from nervous excitement at the beginning of our interview, Marielle stammers as she tells me the story, her scramble for words echoing the anxiety she felt that day when she couldn’t breathe.

‘Well I thought that I was gonna die because it was my first time. It was my first time that I got rheumatic fever?’

I go to clarify, ‘you thought you were going to die-’

‘-Yeah, because it was my first time.’

‘Because it was your first time. How come you thought you were going to die?’ I wonder.

‘Oh because I couldn’t breathe properly?’ Marielle explains. ‘And because like- like if you have rheumatic fever, like, you have to like-’ She gasps, demonstrating. ‘Like breathe like in, and then you have to like, breathe out. It’s like really weird. For the first time, but when you get your second, like, you’ll know.’

Marielle’s breathlessness began when the doctor said she had rheumatic fever, and back at home she found herself processing what that meant. She knew what rheumatic fever was, because the television ‘said it’, even though she did not know how it would feel before. Voice shaking, she tells me, ‘then I was thinking, like I was gonna be that boy?’ Marielle’s retelling of her strep throat experience bristles with embodied anxiety, mirroring the gasping breaths which crept upon her as she went about her day. For Marielle, the experience was ‘making real’ the story she’d heard on the television, and, thinking that she too would be like ‘that boy’, she stressed about taking her pills. Her sister’s wedding was scheduled before the end of her antibiotics course, so to make sure she would be well for the event, Marielle took the final few days’ worth all at once. When I ask her what she remembers from the television
advertisement, she is vague on the details, but she remembers the death part. ‘It was just about how, like when he was born his heart was like, not- good? Then he got a surgery. Something. ‘Cause his brother almost died.’

This is where children’s concepts of death matter. Adults may forget that an audience is wider than the intended target, that adult talk of death will reach children, and that children will make their own sense of what they see. Because death is such a powerful concept in children’s worlds, in linking childhood and death adults inadvertently speak children’s language directly to them.

For these children, rheumatic fever’s representation in media campaigns discursively links illness to death, but also structures health care in terms of death management. The particular version of death which constructs narratives of rheumatic fever also drives children’s practices: their attendance to the clinic, the way they take medication, even their breathing.

Many children at Tūrama School expressed that they would come to the clinic to have their throats checked ‘in case’ out of fear of death. As described in chapter four, Dandre, afraid of the bug that will kill you, goes to the clinic so Whaea Allison can check if he has a sore throat. His sister Jordyn, associates forgetting her medication with a near death experience, and, panicking that she might die, takes her antibiotics ‘again and again’, before she puts the sticker on her chart to confirm the medicine has been taken, and she can relax once more. Marielle conscientiously takes all of her pills to be sure she is cured by the time of her sister’s wedding.

Such hyper-vigilance is again mirrored in Hunleth’s (2017) Zambian children, where fear of ‘orphanhood’ drove children to be exceptionally vigilant in maintaining the health of their parent, while adults endeavoured to remove the child from proximity of the ill parent. Children would respond to their vulnerability by taking responsibility for the monitoring of their family members’ treatment regimes, including ensuring their relative adhered to the treatment by bringing them their pills and drinking water and encouraging them to take their medicine, and becoming quite distressed when hospitals prevented them from monitoring treatment adherence. However, at Tūrama School, clinic nurses also identified the opposite issue, where sometimes children with badly infected throats would avoid the clinic out of fear. Whaea Allison recounted a story of going to visit the home of a child with repeated positive strep A results to swab their siblings in an effort to stamp out the bacteria. A seven-year-old brother with a badly infected throat refused to be swabbed because he’d seen the advertising and didn’t want to get rheumatic fever. Thus, hyper-avoidance, along with hyper-
vigilance, could turn into a health care strategy in the management of death. In this way, children’s concepts of death, constituted as a mode of expression in peer cultures and shaped by the structures of childhood in socio-political context, are woven into practices, and hence into the coproduction of health itself.

Conclusion

This chapter describes one account of how death appears in children’s everyday, in a context where death is both culturally salient and demographically more frequent. While previous anthropological literature dealing with child perspectives on death have primarily considered death as an anomalous childhood event, children in many contexts have extensive contact with death, if not through personal experience, then through the experiences of peers or representations of death in stories, media and games. Listening for death in everyday small talk can hear how children come to not only express their sense-making of death, but how they use death to make sense of life. As an expressive tool, death can therefore also function as a lens for understanding the experience of childhood.

The experience of childhood that Tūrama School children’s death talk reveals is one of embeddedness in relations of kinship and care, and of a dependency which creates precarious social positions. For children, death may invoke not only moments of existential disconcertion, but the social disruption that follows the loss of a parent figure. Although for many, close whānau ties mean that a child who loses one or both parents will often be rehomed with other relatives, children’s dependence and lack of power leaves them structurally vulnerable to unpredictable dislocation. Death is the slipping of your feet on a ground that has suddenly given way, with no promise of where you might land.

From this structural vulnerability, their social dependency and lack of power, death, then, carries special meaning for children. Children’s expressions of death, in particular a process of apprehending this unknowable threat though a social cataloguing of death events and causes, can bring into relief their experiences of vulnerability. Furthermore, the way that children use death language to express ideas about other things in their everyday discourse can also reveal the social meaning of death, and of those things. For children, the language of death comes to connote what death represents as well as its literal meaning, a shared signifier of the scariest experience, the most dangerous risks, or the worst illness. Yet the same structures which render children vulnerable also make them invisible to policymakers who
invoke child death to persuade adults but inadvertently tap into this powerful discourse of childhood, influencing children’s practices in life.
Chapter Nine

Conclusion

In May 2016, six months after I finished fieldwork, I visited Tūrama School again for the day. At the beginning of morning tea I was swamped with excited children who remembered me, while new faces stared with curiosity and suspicion. After five minutes they had, for the most part, run out of things to tell me, and drifted back to their games.

There were many faces missing. Victor, who had the rheumatic heart condition, had not returned to Tūrama School this year, I thought perhaps because his mother Adrienne had decided to move him to a school closer to home. Teuila, who begged spinach from my sandwich and shared her antibiotics, had also not returned this year and a teacher suggested she would now be attending a school closer to the state house her family had been allocated after the death of her father. Nga-Atawhainga and her sister Te Rerenga, Harrison, Amberlee and Arya had all moved away. Te Kapua, Tupono, best friends Trystan and Navahn, and many others, having completed year six, had graduated to the intermediate school nearby. Some of the staff had changed as well. Deb, the nurse, was still working in the clinic, but Whaea Allison had moved on to another role and there was a new ‘sore throat lady’ now. I had arranged the visit with Mrs Randall, but when I arrived found she was sick and had not come to school that day. I was not to know it then, but within a few months she would take an early retirement due to illness, and I would not see her in a classroom again.

But I was delighted to find Cassidee, who had told me last year she thought she would be moving again, now in the year 7/8 bilingual unit. She still had her stoic poise, but had paired up with Soraya, who used to be best friends with Ruby, and both girls had matching shaved undercuts, the popular hairstyle. They saw Ruby from time to time, they told me, when she came to the gate to meet her little sisters, but she didn’t really speak to them. I also spent some time with Mila, who was growing into a leadership role in Mrs Steven’s class this year. She, Hinemaia, and Dandre all proudly sported school councillor badges. Pikau, now year eight, has grown tall and lanky, well into adolescence.

Around the time that I began writing this conclusion in 2017, a UNICEF report on child wellbeing in developed nations had just been released, ranking New Zealand 38 out of 40 in a measure of health and wellbeing across five indicators: neonatal mortality (<4 weeks), suicide rates (0-19 years), mental health symptoms (11-15 years), drunkenness (11-15 years)
and teenage fertility rates (15-19 years) (UNICEF Office of Research 2017). At a rate of 15.8 per 100,000, New Zealand’s youth suicide rate is the highest of any developed nation by a substantial margin. These are the issues awaiting the children of this thesis; as their immune systems mature and they grow out of vulnerability to rheumatic fever, they age into new demographics of risk. In the most recent report from the Ministry of Health, the rate of youth suicide in Manukau, the district health board that includes Papakura, is higher than the national average (Ministry of Health 2016). The same report shows suicide rates increasing with deprivation quintiles; a relationship strongest for the youth population where the three most deprived quintiles experienced four times the number of suicides than the least deprived. Māori and Pasifika young men between 15-25 are the two most at-risk groups, followed by Māori women and European men in the same age-bracket. Highest rates of hospitalisations for intentional self-harm were recorded for young Māori and Pākehā women in the 15-24 age group living in neighbourhoods of high deprivation (Ministry of Health 2016). It seems that the environments many of our young people are maturing in are not even close to providing the resources for thriving.

In conservation science, sentinel species are organisms that are particularly sensitive or manifest early responses to environmental change, and thus are used as bio-monitors, signaling early warnings of contaminants or declining ecological systems. Across time and space, different indicators may emerge within particular sectors of society which represent ‘sentinel diseases’: a canary in the coal mine for inequitable conditions, or a warning sign of deeper faults in housing and socio-economic circumstances that will also manifest in a broad range of other health issues. Such indicators—diabetes, cardiovascular disease meningococcal B, low birth weight, and currently in New Zealand, rheumatic fever—may not be the earliest signs of deteriorating environments, but their measurability and reliable connection to environmental circumstances mean they can function as proxies for material inequities, racism, or spiritual dislocation (Walters et al. 2011).

Yet because treating the canary with antibiotics can show quantifiable effect, attempted treatment of isolated diseases through ‘targeted’ approaches to public health may be preferred not only for financial reasons, but for governments keen to demonstrate progress to those to whom they are accountable. As a Better Public Services (BPS) target of the National Party government, the goal of the Rheumatic Fever Prevention Programme was to cut rheumatic fever admissions by two-thirds by June 2017. As of December 2016, this programme has seen moderate success, with a 23% reduction in rheumatic fever cases overall—in real terms,
from 177 cases in 2012 to 137 in 2016 (Ministry of Health 2017). Although this includes a 42% drop in the Counties Manukau district, where Tūrama School is located, this district still has the highest rate nationally by a relatively large margin (7.9 cases per 100,000 compared to a national average rate of 2.5 per 100,000). Nationwide statistics show a steady decline for Māori rates, but these high Auckland rates particularly reflect the Pasifika statistics, which, after dipping in 2014/2015, spiked again in late 2015 and in 2016 reach the same levels as in 2012 (Figure 11). For all the targeted interventions, Pasifika peoples have seen little change.

![Figure 11. Rheumatic fever rates in New Zealand from 2009-2016. Data source: Ministry of Health, 2017](image)

The 23% decrease nationwide falls well below the targeted reduction of 67%, but was enough for the government to claim success and commit funding for another five years to rheumatic fever prevention. The problem here of course, is that while children have been fortified with antibiotics, the conditions that created the epidemic remain in place. Scaling back the interventions would leave children at risk again, and removal of the school clinics would leave a cohort of children, attuned to sore throat risk, conditioned to seek treatment, with no place to go.
New directions in child policy

Yet the political milieu that seemed so entrenched and stagnant just a few months ago now anticipates a potentially significant new direction. An unexpected change of government in late 2017 through a coalition agreement has seen a rapid reprioritising, at least in rhetoric, of child poverty issues in Aotearoa. An early move of the new, left-leaning government has been to scrap the BPS targets of which reducing rheumatic fever rates represented one (unmet) target. Instead of targeting streptococcus infections, the government has announced it will be focusing on systemic change—addressing the overcrowding and child poverty that underlie rheumatic fever rates (The New Zealand Government 2018). Although exactly what this will look like remains unclear, it appears that the New Zealand government’s approach of medicating the canary may be shifting towards changing the coal mine.

Meanwhile, the outgoing National party, who until the 2017 election disputed both the number of children in poverty and the measurability of child poverty (Moir 2016), criticised the new Labour government’s child poverty reduction proposals as under-ambitious, and, now in opposition after nine years in power, has signalled that it will only support the Government’s proposed legislation if amendments are made to reintroduce BPS targets and ‘hold the public service accountable’ (New Zealand National Party 2018). Hon. Paula Bennett, who as Minister of Social Development rejected legislation to provide children in poverty with free lunch on the grounds of ‘parental responsibility’ (see chapter five), now states her expectation as the National Party Spokesperson for Children that the government would ‘address the drivers of social dysfunction’ and use ‘social investment models’ so that families and communities get the resources they need. Children are therefore very much implicated in current contestations for political power, though it remains to be seen whether this will result in any material change for the incoming cohorts of Tūrama School children. Nevertheless, while Aotearoa still trails countries such as Sweden for child-centred policy (D’Souza 2017) there appears to be fresh reconsideration of the role the state plays—whether through action or inaction—in shaping children’s health and welfare.

Without understating the significant impact of the state and society on children’s health, in this thesis the case of rheumatic fever prevention has been the entry point for a different set of questions, this time about the role children themselves play in the production of their health. Children may be increasingly the focus of policy change but still missing is a view of children as social actors, whose experiences, understandings, interpretations and practices
matter and can powerfully mediate the success of failure of policy interventions. Instead, the trend in New Zealand, as well as internationally, has been towards what scholars describe as an ‘elevation’ of children within neoliberal policy, with the purpose of targeting ‘vulnerable’ children for intervention within a social-investment paradigm concerned with reducing future costs to the state (Elizabeth and Larner 2009; Keddell 2018; O’Brien 2016). The rheumatic fever prevention programme is a classic example here, along with recent changes to child protection policy which channel resources into identifying and policing at risk families rather than providing support or addressing chronic entrenched poverty, material deprivation and mental health issues (Cleland 2016; Keddell 2018).

This, therefore, is ‘child-focussed’ policy, but it is rarely child-centred, in that the agenda is designed for the benefit of the taxpayer rather than for the child, legislation which punishes families for poverty is not particularly helpful for children, and impacts on children are not considered from their perspectives. Other examples of this tendency to overlook the importance of children’s views can also be seen in recent changes to family law which eliminated the rights of children to have a lawyer representing their views in family court unless there are concerns for the child’s safety, in the Adoption Act 1955 which has no requirement for the court to hear from the child (Cleland 2013), and in the rebranding of Child Youth and Family Services to the stigmatising and controversial ‘Ministry for Vulnerable Children’ (though another early move of the incoming government was to remove the word ‘vulnerable’ from this name).

Fitzmaurice (2017) notes that Aotearoa lacks the participation ecosystem—the participatory culture, infrastructure, understanding and research—needed to allow children’s perspectives to shape institutions, practices and services in useful and meaningful ways for children. When children’s perspectives are included, such as in the recent consultative processes for the reformed child protection services (Fitzmaurice 2017), these still tend to remain adult-centric in process, transplanting children into adult political arenas (Lansdown 2010), and with an emphasis on voice, meaning the verbal sharing of children’s experiences, views, and opinions, which privileges the views of older and more articulate children. While children can certainly express rich and valuable perspectives, as many did in the present study, a child-centred approach to policy must also hear children’s many non-verbal expressions and situate these in the ecological, relational, cultural and institutional contexts within which they have been produced. This requires knowing children’s lives, embodied, negotiated, and embedded in relationships, beyond what children might name in an interview in a strange
place with strange adults. The ethnographic mode of ‘being there’ (Borneman and Hammoudi 2009) is particularly useful for understanding the perspectives of children who may be less practised at verbal articulation but who can nonetheless express their experiences through multiple other, embodied or relational modes. Such an understanding of children’s lives would better allow the systematic and reflexive suspension of adult-centric lenses and dominant social conceptions of children needed to instead view processes and policies from the child’s eye view. This also allows consideration of potential effects of policy decisions on children’s practices, whether that is the way a stigmatising Ministry name may affect children’s peer relationships, or the way a rheumatic fever campaign may create hyper-vigilant health practices.

How do children participate in the coproduction of their health?

How, then, should we conceptualise children’s contributions to their health in the context of their lives? While researchers embrace questions of how children experience health and illness, their interpretations and practices rarely figure into analyses of the production of child health. In New Zealand such analyses take the form of mainly longitudinal quantitative studies (Fergusson, Boden, and Horwood 2015; Savila et al. 2011; Morton et al. 2013; Silva 1990), where, perhaps, children are presumed passive, or their activities disregarded as inconsequential, or their social action assumed to be the product of their environment and what adults are doing. Some of the reticence to seriously consider children’s practices may stem from an idea that children ought not be consequential players in their health, that children’s health should be produced by adults, or at the very least, children’s role should be learning to take care of themselves as they are taught at home and school. To focus on children’s contributions to their health status risks making them responsible for their wellbeing, or shifts attention from the many ways that their health is influenced by people and social forces with far greater power. Yet how would our understanding of child health change if we were to include a view of children’s interpretive practices as part of the analysis?

I have argued throughout this thesis that such a view of children’s activities would reveal that children’s practices, generated through collective peer cultures and in relation to wider social structures, have significant potential to structure their health. I have shown how children’s interpretive practices can shape their nutritional status, use of pharmaceuticals, engagements
with health care, allostatic adjustments of the body’s internal milieu, social support and exposure to risk. These practices can also influence the success or failure of policy or institutional interventions intended to facilitate their wellbeing. These are not particularly new findings; similar sorts of practices can be found throughout the existing interpretive literature on children’s health and illness: how children employ imaginative strategies to cope with chronic illness (Clark 2003); how children interpret their bodies in culturally specific ways and consequently may under or over-report symptoms (Christensen 1999); how children within peer cultures stigmatise some forms of illness (runny noses, eczema) (James 1993); how children attempt to stay in proximity to and care for an ill parent (Hunleth 2017). There are clear implications of these practices for children’s own health; we might anticipate they mediate treatment adherence or efficacy, timing or accuracy of diagnosis, social support and wellbeing, or exposure to pathogens. These are not inconsequential effects, but in this literature the conversation is not about how these activities help to make child health.

How, then, can children’s practices be integrated into conversations about the production of child health? How can we represent children’s agency in analyses of rheumatic fever, obesity, or depression without implying children are responsible for their health status, making their behaviours into targets of intervention, or detracting attention from structural inequities and injustices? One solution has been offered by the social ecology models of childhood resilience (Panter-Brick 2015; Ungar 2011; Ungar, Ghazinour, and Richter 2013), discussed in chapter seven, which attempt to include children’s as social actors who negotiate their own wellbeing in relation to environments which constrain or open up resources to facilitate young people’s practices. This sets up a way of giving attention to children’s practices as strategic choices made from a limited number of available options, and in view of children’s circumstances, practices which may seem counterproductive from adult perspectives can be understood as accommodations which balance costs and benefits across different domains in resource-poor or challenging contexts. A view of what children do is valued here for understanding how to improve environments and resources in ways that will give children better options or mitigate the costs of what children do.

Working from psychology and social work fields (though also picked up in the large-scale anthropological work of Catherine Panter-Brick (e.g. Panter-Brick et al. 2015)) the social ecology model tends to take an individualised and unidirectional view of children responding to the conditions of their environment. What the ethnographic data from Tūrama School shows, however, is the way that children’s practices are deeply interconnected, and, as such,
work to help create the environments that constrain and open up opportunities and resources. Children like Ruby work within peer ecologies to create friendship groups that stabilise an unstable world, but at the same time, Ruby’s practices create new vulnerabilities for the children she excludes. Boys like Harrison, who employ aggression to maintain their social safety, create unsafe environments for their targets. Children technically have access to spare lunches, but the social meanings they collectively ascribe to these mean that this form of resource is constrained by social costs, while other forms of provisions, like the milk, are woven into collective, and enabling social rituals. This dialectic, where children collectively co-construct the structures which guide their action, is not quite captured in existing models.

I have therefore argued that a coproduction framework offers another way for us to make room for children’s agency by positioning children as working in dialectical relationship with adults, the body, and wider social structures including the state, the institution, cultural ideologies, and the economy. Coproduction, as I have used the term—the way that children, society, and the body make each other—recognises that children do not simply internalise, but creatively produce culture (Corsaro 1992). Yet these cultural meanings and practices, on the one hand, are structured by experiences of the body and wider social systems, and on the other, mediate the relationships between wider social systems and the body; they are coproducing and coproduced. Thus, Tūrama School children construct understandings of sore throat at the interface between their experience of the body incarnate and a political agenda that targets sore throats within health care systems (chapter four), and interpret provisions as acceptable or stigmatised care through their own notions of intergenerational roles and responsibilities (chapter five). From their experience of structural vulnerability, they make meanings of death in terms of illness and threat to the social (chapter eight). Meanwhile, these cultural meanings themselves form structuring social systems that guide children’s practices and become embodied as a particular form of habitus (Bourdieu 1984), directly shaping their health. Peer cultures create systems that constrain food resources or open up access to health care, which then structure how bodily signals of hunger, cold or sore throat are experienced, interpreted, and responded to (chapter six). Children’s understandings of social relations, responsibility for care, illness, medicine, and their own bodily sensations influence how they engage with health care, patterns of nutrition, or practices of using pharmaceuticals (chapters 4-5). Children strategically employ practices of resilience to enable coping in one domain of health (e.g. peer society) but these may come at a cost in another (e.g. risk of physical injury) (chapter seven). Through these processes children translate their
politically and economically structured experiences of the world and of their body into local social meanings and practices, which in turn help to shape future action and future bodies. Children’s health activities—building relationships, caring for their body, eating and drinking, taking medicine, engaging in risk—are all coproductions authored by society, families, institutions, and children themselves. Children are not solely the ones responsible for their health here, but institutions and policy-makers are made responsible for considering how children will understand and respond to the conditions of their childhood, and for taking children’s meanings and practices into account designing or evaluating policy or social services.

Viewing children’s health as a coproduction enables us to value and give attention to children’s efforts to manage their own lives and support the health of others. In practice, as well, acknowledging the role children play within these dynamics allows us to better predict the effectiveness of policy decisions, or better still, design interventions and services that align with and support children’s ideas and goals. Such an approach might ask: what would school lunches look like if created with a view of how food functions within children’s social relations? How would rheumatic fever policy change if we began by asking what a sore throat means to children? What if child protection began not with how we can identify, monitor, and police ‘at-risk’ families, but how can we support and preserve those aspects of families that children value and are trying to make for themselves? There may be a near consensus that poverty is bad for children, but when viewed through children’s eyes, some responses to child poverty are also bad for children. From children’s perspectives, material resources may matter less than the effects those resources have on family relations, time, and affective care. What kinds of resources, and delivered in what way, would make the biggest difference to what children need and value?

A coproduction framework also allows us to better understand how the systems that generate health inequalities operate through childhood experience, and how society discursively and structurally positions children and influences their activities. When viewed as coproductions, children’s perspectives and practices tell us something about the way society is structured to create ethnic, economic, or generational vulnerabilities and how these are experienced. The way children use the language of death in association with illness speaks to the precariousness with which they view their life, and how health inequities destabilise their social relationships and sharpen their perceptions of risk. A coproduction framework may
also be applied outside of health, as an analytical tool that moves beyond socialisation or internalisation for understanding how children contribute to other aspects of their childhoods.

Finally, viewing children’s activities as a coproduction also means considering the ways that children contribute to their coproduction partners: how children help to support adult health, how children modify institutions, how children influence society. Children’s lack of social or political power can sometimes be mistaken for a lack of effect, however, children shape economies as consumers (Buckingham 2011) as well as labourers (Kramer 2005; Nag et al. 1978), act as language brokers for immigrant parents (Orellana 2009), and co-construct learning exchanges with grandparents (Kenner et al. 2007). Jean Hunleth’s (2017) work on children’s roles in caring for parents with tuberculosis—what could be termed the coproduction of family health—argues for the importance of not limiting consideration of children to ‘children’s issues’, or only those activists, researchers and policy-makers concerned with children. In Hunleth’s case, the invisibility of children from the perspectives of health professionals caring for adults, and attempts to remove children from adult tuberculosis sufferers were in themselves harmful to children. Likewise, at Tūrama School, the invisibility of children to rheumatic fever campaign designers created anxious children. As illustrated by Mila’s story in chapter seven, the activities and positions of children and adults are intertwined, so just as parents fret over taking their child with a sore throat to the doctor, children will worry about a parent who frets. Social discourse therefore cannot speak of parents without speaking to children, and vice versa. A coproduction framework can remind us to look for children’s influences, as well as how they are impacted, in studies of adults or society more broadly.

This thesis has been about elevating the unseen and underestimated practices of childhood in the social theory of health and of society more broadly. I have also spoken to how we should consider childhood in practice: of grounding policy in understandings of children’s experiences and activities as they are lived in situ. Most importantly for me, this thesis has been about listening to, and hearing from a group of children who feature regularly in public discourse as statistics and stereotypes, but whose stories as people are rarely heard. The word ‘tūrama’ means light, and I named Tūrama School for the shining little humans I saw every day. Many of these children suffer greatly the effects of structural violence, but their lives are also of helping friends and joking with teachers, of sneaking lollies to share, of loving their parents and grandparents, of taking pride in their mathematics and soccer and art, of caring,
grieving, singing, fighting, resisting and trying. These are the practices of childhood, and they are the most bright and brilliant things that I have ever seen.
Appendix

Picture book produced for child research participants, teachers and school.
In 2015 an anthropologist came to our school to learn from us about our lives.

She talked to us and drew pictures of what she saw, and we drew pictures for her as well.

This is the story of what we taught her.
We are kids.
We live in a world built by adults,
but we make it our own.
(The toilets are an important place for secrets).

We create worlds of meaning that adults can’t see.
We like to talk, but much of our language is unspoken.
Swapping shoes shows best friends.

Doing dares shows you belong in the group.
Hugs say I care for you... or I want you to care for me.

We have our own etiquette for play.
If you want to join in, you must first ask the boss of the game (usually the person who started it).
But it is polite for the leader to let others join.
It hurts when they say we can’t join.
We love to practise our skills on the playground: on the spinners, on the jumping logs, skipping ropes and gymnastic bars. We show off our mastery.

Lots of us love soccer.

We are the best players in the country! (And we came third in the local tournament).

We can be flexible with rules, especially when it helps us win!

But we have a natural sense of equity too.

We’ll pretend we don’t see when a new player does a handball.

School can be a hard place to be sometimes. But when we are with a friend, school feels much safer.

Our friendships are protective, and so vital to our resilience.

Our friends are the people who have fun with us, care for us, keep us safe from bullies.
So adults, please understand:
When we have a fight with our friend...
When our friend leaves us for someone else...
When our friend teases us and makes us cry...

When our friend moves away...

This is a big deal.
We will always remember the way it affected us.

They take care of us when we are sick.

Our whānau are very important to us.
If our friend is our cousin, then that is a doubly special relationship because blood doesn’t change.

So we claim our whānau ties,

And call our friend our brother to make him more important.

We live in a world where people are always changing.

Friends leave, and new kids arrive...

Parents and relatives come and go.

People die.

And we usually don’t have much say, or understand what’s going on. (And that can be scary.)
Being a kid is pretty awesome.
We don't have many responsibilities.
We don't have to have a job,
or look after children.
We can eat lollies and play.
We are free.

Adults have to clean.
Adults must pay bills.
Adults have to work.
Adults fight.
Adults can drink and smoke.
Adults forget things about what
it's like to be a kid.

Sometimes we are scared that when
we're adults we might do things we regret.
It's not always easy being a kid.

But we're pretty sure adults wish they were us.

So we reckon we're the lucky ones.
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