Does your carer take sugar?


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Luke was involved in the drafting of the Bill’s that became the Carers (Recognition and Services) Act 1995 and the Carers (Equal Opportunities) Act 2004, as well as acting as the expert adviser to the Westminster Parliament’s Joint Committee scrutinising the draft Care & Support Bill (2013) and the Welsh Affairs Select Committee on the Welsh Government’s Carers Strategies (Wales) Measure 2010.

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Does your carer take sugar?


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Abstract:

This paper considers the struggle being waged by unpaid carers (sometimes referred to as ‘caregivers’) for recognition as ‘rights holders’. It locates the origins and describes the growth of the ‘carers movement’ and argues that it has many similarities with the Disabled People’s movement that came to prominence in the 1970’s. The paper: (1) identifies the distinct legal status that carers have in the majority of states in the world; (2) describes carers’ shared history of adverse treatment within most states; and (3) argues that carers’ social exclusion arises from a widespread hostility to ‘dependency’ – a hostility that is gendered and particularly evident in neoliberal political discourse.

The paper argues that there is a substantive human right ‘to care’ – one that fits most comfortably within the civil and political right to ‘privacy / private life’; that states have positive human rights obligations to carers; and that ‘being a carer’ should (and will) become a protected status for the purposes of non-discrimination legislation, on the same basis as other protected statuses (such as disability).

The paper concludes with a caveat: that the recognition of caring as a human right and of carers as ‘right holders’ (although inevitable and of great importance) will not in itself be sufficient – that this much we also learn from the Disabled People’s movement.

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Does your carer take sugar?


35 years ago a BBC Radio programme came on air in the UK called ‘Does he take sugar?’

The title was provocative, since the subject matter of the programme was disability. The title tilted at what it perceived (rightly no doubt) as the prevalent conceptualisation of a disabled person by its listeners: that of a compliant cripple seated in his wheelchair grateful for the services and sympathy bestowed upon him. A tragic unhearing victim, incapable of expressing independent opinions or knowing what he needed – an object of discussion, whose needs were primarily the responsibility of the social care authorities.

The programme was a symptomatic marker of the journey disabled people were travelling: towards a radical transformation of the way they understood themselves and the way non-disabled people understood disability. At the time of the programme a few legal milestones had been planted, of which the USA’s Rehabilitation Act 1973 was a prominent, if modest example.

Today the popular conceptualisation of a disabled person would, I hazard, be quite different. The tragedy module still no doubt dominates – but the notion that disabled people are compliant, grateful and non-verbal would be discounted by the vast majority. To use the phrase ‘a disabled person’ is to conjure up the associated notion of discrimination legislation: of a group who are potentially prickly – who sue and who take direct action if you get on the wrong side of them. They are conceptualised by an increasingly large portion of the population as rights holders: people who win human rights cases and for whom it is no longer the social care authorities (or even public bodies) who have sole responsibility: all of us now have responsibilities – schools, cinemas, supermarkets, banks – even Ryanair.

In the mid-1970’s few lawyers would have considered disabled people as candidates for non-discrimination legislative protection. Sex and race maybe (contested as of course these had been) but disability was such an elusive concept, and in any event it was viewed as a self evident handicap, unlike sex and race which were (by then) viewed as prejudicial inferiority constructs: lacking any material justification. Disabled people on the other hand, needed state supports and accordingly were better conceptualised in terms of positive obligations and as having the protection of the soft socio-economic rights rather than the hard negative civil and political rights.

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With the widespread acknowledgment that disabled people are materially handicapped by social and physical barriers (the so-called ‘social model’ of disability) such an analysis is no longer tenable. In individual domestic legislative terms this new conceptualisation is manifest in the surge of provisions outlawing disability discrimination,3 which in turn led to regional and international action culminating in the UN Convention on the Rights of Persons with Disabilities. In little over 30 years we have redefined ‘handicap’ and have come to regard it as normative to view disabled people as entitled to equal treatment: even if the prevalent conception of a disabled person is still a ‘he in a wheelchair’.

Today when human rights lawyers consider the language of the key founding documents, for example the International Bill of Human Rights, the American Convention on Human Rights and the European Convention on Human Rights, many express surprise at the absence of disability from the familiar litany of protected statuses: race, colour, sex4 – as if disability was a self evident category for protection: indeed it was not.

A while ago I wrote to the BBC to suggest that they should broadcast a new programme: ‘Does your carer take sugar?’ – for it is at least arguable that carers (by which I mean people who provide care on an unpaid basis for a ‘dependant person’5) find themselves in a position similar to that of disabled people 35 years ago. Today the prevalent conceptualisation of a carer is, I would suggest, of someone grateful for the services and sympathy6 bestowed upon her, and for whom the social care authorities have prime responsibility. Few human rights lawyers would conceptualise carers as rights holders: for being a ‘carer’ is an elusive concept and in any event it is not an innate characteristic (like sex, race and disability) and the handicaps experienced by carers are those they assume when they choose to take on their caring roles: carers’ need for support is therefore better conceptualised in terms of positive obligations and as having the protection of the soft socio-economic rights rather than the hard negative civil and political rights. Carers, in a phrase, cannot legitimately be viewed as ‘rights holders’.


4 See for example, Article 14 European Convention on Human Rights which specifies ‘on any ground such as sex, race, colour, language, religion, political or other opinion, national or social origin, association with a national minority, property, birth or other status.’

5 Generally referred to as caregivers in the USA.

In this paper I argue that carers should be seen as a category of persons entitled to protection from discrimination: that they should and will come to be viewed as self evident ‘rights holders’. By contrasting their two journeys I do not argue that they must follow the same route: all discriminations are unique and although there are profound similarities between the struggles waged by people subjected to discrimination on grounds of sex, race, disability, sexual orientation, age, religion and so on – each of these ‘statuses’ has its own inimitable core and distinct narrative. There is of course an obvious interconnection between the struggles of carers and disabled people for equal treatment, but this connectedness should not mask the challenging differences between their claims for recognition.

The Rights Moment

For a group experiencing oppression to unite and to express their disadvantage in the language of ‘rights’, depends upon the convergence of a number of socio-political, cultural and conceptual factors. This paper considers three: the development of an identity; a narrative; and a creed.

The group needs, in one way or another, to ‘self-identify’ as a category of persons oppressed by virtue of a particular uniting characteristic: as Shakespeare has observed7 (in the context of the development of the disabled person’s movement) an identity that ‘connects the social and the personal and involves the individual putting themselves in a collective context’: a context that ‘focuses on… exclusion and injustice’. It is a process that needs a history: a narrative documenting the nature and the extent of the negative treatment they have experienced. Finally the group needs a convincing theoretical model that articulates and explains their adverse treatment in social and political terms. For disabled people, of course, this was the social model of disability.

Once these factors are in play, there is the potential for a radicalised campaign, challenging all aspects of the negative treatment experienced by the group: one that demands equal treatment in place of toleration or ‘privileges’. It is the stage at which socio-political and economic forces converge creating a ‘constitutional moment’8 - an imperative for legal change.

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Identification

As a simple matter of human rights and equality law, discrimination becomes unlawful when a person is treated unfavourably for a ‘status’ related reason – provided it is a recognised or ‘protected’ status.

The legal bestowal of ‘status’ is an explicit and highly symbolic act: evidence that society attaches such importance to a distinction that it demands explicit recognition. It is the stuff of power relationships and tribalism: of critical masses and crystallisations. It involves the assigning of a value to a difference: though in truth the process has generally the effect of devaluing – whether articulated in the language of paternalism (as it has been for women and disabled people) or the language of criminalisation (as with Gay or Aboriginal / First Nations Peoples).

Whilst the legal recognition of status is a necessary precursor to any rights movement – it is, in itself, insufficient. The group has to take possession of that status, redefine and own it. Though the process of reconceptualisation may often require a renaming (no longer nigger, cripple or queer) it will invariably be subversive and celebratory: of Ian Dury, Gay Pride and sisterhood – for the law and social change, like shackled prisoners, move together. The group must, in short, ‘self-identify’: its members must embrace sabotage and radicalise the status it has been assigned.

The section that follows considers these two issues. It commences with an overview of domestic laws that deal with carers as a specific group. It then considers the socio-economic and political factors that have produced the essential component for legal change – a critical mass of self identifying carers.

The legally entrenched status of carers

In much of the world, ‘being a carer’ is not only a designation that results from a process of self-ascription – it is also a legally created status. Whilst scholars differ as to the reasons for such provisions,9 laws obliging people to provide care, solely by virtue of a family relationship or marriage, are ubiquitous.

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9 It has, for example, been suggested that there are philosophical – rather than simple community cost avoidance reasons for such obligations see for example MC Stuifbergen and JIM Van Delden, ‘Filial obligations to elderly parents: a duty of care?’ Medicine, Health Care and Philosophy (2011) 14: 63
In the UK a liable family rule was formalised in the Poor Relief Act 1601 and persisted (though reformulated in the Poor Law Act 1930, s14) until repealed by the National Assistance Act 1948, s1. The Poor Law was exported to the colonies where it has proved to be more tenacious. In the USA, for example, it appears that 30 states¹⁰ still retain filial responsibility statutes.¹¹ Such laws are also present in most Canadian states¹² (albeit endangered¹³) in India¹⁴ and Singapore.¹⁵

In Europe, duties on family members to provide care (or financial support in lieu of care) are found in the Constitutions of Greece and Ireland¹⁶ and in the Civil Codes of many states – for example, Belgium,¹⁷ France,¹⁸ Germany,¹⁹ Italy²⁰ and Spain.²¹ The Civil Codes of many South American Constitutions contain a similar obligation (the duty on family members to provide ‘alimentos’) for example in Argentina,²² Peru²³ and Brazil.²⁴

¹⁰ Alaska, Arkansas, California, Connecticut, Delaware, Georgia, Idaho, Indiana, Iowa, Kentucky, Louisiana, Maryland, Massachusetts, Mississippi, Montana, Nebraska, New Hampshire, New Jersey, North Carolina, North Dakota, Ohio, Oregon, Pennsylvania, Rhode Island, South Dakota, Tennessee, Utah, Vermont, Virginia, and West Virginia
¹² See for example, British Columbia’s Family Relations Act 1996 s90 and Newson v Newson, 99 BCLR 2d 197 (1994, BCSC); Saskatchewan’s Parents’ Maintenance Act 1978, s2; Manitoba’s Parents’ Maintenance 1996, s1 and Ontario’s Family Law Act 1990, s32
¹⁷ Articles 205 - 207.
¹⁸ Articles 205 and 206
¹⁹ Bürgerliches Gesetzbuch (para 1601 Civil Code) entitles close relatives to financial support against each other in times of need although this can be financial, as opposed to the actual provision of social care – see Meyer, M (2004) National Background Report for Germany EUROFAMCARE Hamburg para 2.1.4; see also Means & Smith, ibid p220.
²² Civil Code Articles 367, 372 and 376 (bis).
²³ Articles 472 and 474.
²⁴ Articles 1694 and 1696.
The obligation is found as far afield as in the Civil Code of Taiwan\textsuperscript{25} and in the customary laws of some African states.\textsuperscript{26} In such customary laws (as indeed in the Irish Constitution) the gendered nature of the obligation is explicit: it is the duty of wives / women

Even in those states where such obligations do not exist, or where the obligation is not in practice enforced, the evidence suggests that the dominant social attitudes exert strong moral pressure on family carers to fulfil this role. The assumption being that the family has primary responsibility for care giving: indeed, not so much family, as women whose duty it is to look after sick and frail elderly parents and in-laws.\textsuperscript{27}

In Australia, where no filial responsibility laws exist\textsuperscript{28} it is said that there is an ‘expectation that families will take the primary role in looking after the elderly members’\textsuperscript{29} and in the Netherlands although there is no formal legal duty to provide care, the assessment of need under the state’s Long Term Care Insurance Scheme includes an amount of ‘customary’ care family members are expected to provide for each other free of charge.\textsuperscript{30}

In much of Asia, it is said that the ‘Confucian ideal of filial piety is ubiquitous’\textsuperscript{31} and to be highly gendered:\textsuperscript{32} in Japan for example these values create the assumption that ‘middle-aged women’ will provide the home nursing required by infirm elderly relations.\textsuperscript{33}

\textsuperscript{25} Article 1114 of the 1929 Civil Code.

\textsuperscript{26} Ghana’s customary laws, for example, make it the duty of the wife and children to support their husband and father – see Ollenu J. (as he then was) in Quartey v. Martey & Another (1959) GLR 377 and E Dankwa, ‘Property Rights of Widows in their Deceased Husband’s Estate’in [1982-85] 16 University of Ghana Law Journal 1, 2.

\textsuperscript{27} For the UK context – see Means, R. Richards, S. & Smith, R (2008) Community Care: Policy and Practice (Public Policy & Politics) 4\textsuperscript{th} ed Palgrave Macmillan p 218.


\textsuperscript{29} S Sutherland The Royal Commission on Long Term Care With Respect to Old Age: Long Term Care - Rights and Responsibilities Cm 4192-I (Stationery Office, (1999) p.201.

\textsuperscript{30} Glendinning, C and Moran, N (2009) Reforming Long-term Care: Recent Lessons from Other Countries Social Policy Research Unit, June 2009 Working Paper No. DHP 2318 para 3.2.2. and see also A Struijs, Informal care: the contribution of family carers and volunteers to long-term care (Council for Public Health and Health Care, the Netherlands, 2006) p 66


\textsuperscript{32} H Zhan R Montgomery Gender and Elder Care in China Gender and Society, Vol. 17, No. 2, (2003), pp. 209-229

In the USA, where few of the filial responsibility statutes are actively enforced, federal policy requires that social care plans detail the ‘expected participation of informal carers’ to ensure a ‘reasonable division between informal and formal support systems’. In similar fashion, in Germany it is suggested that the ‘internalization’ of the traditional family caring role/ responsibility continues to be an important factor for individual caregivers. So too in Ireland where, although the Constitutional obligation is not litigated, there exists a ‘strong moral obligation’ on families to provide care. The position is said to be the same in Spain and Greece where (regardless of the legal situation) women have internalised their role as carers – often with materially adverse impacts on their physical and mental health.

Compensatory provisions

A significant literature exists that critically examines various examples of positive state action to address the needs of carers, particularly where these take the form of direct financial payments. In addition to such arrangements, a wide variety of other ‘carer compensation’ provisions exist in the domestic laws and policies of many states. These may, for example, provide for indirect benefits (eg through the tax or pension assessment systems) or mandate flexible employment rights.

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38 Means and Smith cited above, p221.


'Carer Recognition' statutes exist in Australia and the UK, and specific measures to support carers have been acknowledged as a political priority in the majority of European states with many providing for formal (but generally modest) 'carer' payments or 'respite' care arrangements – Finland, France, Hungary, Spain and the UK for example. In Canada a Compassionate Care Benefit scheme for working carers has been developed as part of the Labour Code. In the USA the National Family Caregiver Support Program provides for grants to states to fund a range of supports designed to sustain the care provided by informal caregivers and specific provisions exist for the caregivers of veterans. Official carer specific measures are not solely a Western manifestation: they can be found in (for example) Taiwan, South Africa and India - and are being proposed in Columbia.
Self identification as carers

The acquisition of a social identity is a distinct process for every marginalized group – albeit that there are general and reoccurring themes. In terms of the struggle for human rights, such self categorisation is invariably bound up with the idea of oppression and of ‘imposition’: of a collective identification with unjust subjugation.58 Shakespeare refers to the particular conceptual difficulties that disabled people had in this respect (compared to ‘women, blacks, or gays’), in that the oppression they experience is ‘couched in terms of paternalistic support and charity’.59 Clearly this observation is particularly apt in relation to many carers. Many carers consider caring to be an inherently private, family and charitable activity: it is very commonly reported that many people providing care in such situations ‘do not identify themselves as carers’.60

Identification based on ‘being a carer’ has the additional complexity for those who perceive that their caring role has robbed them of their (former) status – that like the acquisition of an impairment – it has resulted in a lost sense of self identity.

Many accounts that document the radicalization of disabled people in the 20th Century locate its origins in the USA, with Vietnam veterans returning to experience the handicaps imposed on them by environments constructed by and for non-disabled people. Vietnam created a tipping point, by generating large numbers of young, educated and physically impaired people who experienced adverse treatment for the first time in their lives and for whom the activism of the civil rights movement acted as a catalyst in the development of their group coherence.

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58 The social identification of carers and the extent to which they can, as a group, be viewed as a new social movement lies outside the central purpose of this paper: in this respect, however see Tom Shakespeare, ‘Disabled people’s self-organisation: a new social movement?’ in Disability, Handicap and Society., (1993) 8, 3, pp. 249-264 and in particular his critique as to the extent to which ‘post-materialism’ is a core feature of such movements: see also Alberto Melucci, ‘Challenging Codes: Collective Action in the Information Age’ (Press Syndicate 1996) and John Turner, ‘A Self-Categorization Theory’ in John Turner (ed) Rediscovering the Social Group, A Self-Categorization Theory (Blackwell 1987) pp 18–41.


The exponential growth in disabled people’s organisations in the decades following Vietnam is being mirrored by the remarkable growth in carers’ organisations, alliances, networks and support groups that has occurred in the last two decades. As with disabled people’s groups these come in all varieties — local, user specific (e.g. by the nature of impairment, ethnicity, age, or sex) international and so forth. The recent proliferation of organisations of self-identifying carers’ results from the spectrum of socio-legal factors addressed in this paper — particularly from the impetus created by their recognition in domestic legislation and other formal policies. These measures are, however, merely a response to wider social forces. A number of commentators argue that for carers, the significant motive force the ‘Vietnam’ issue is the impact of neoliberalism — and its disparagement of dependency.

It is however the coincidence of this political phenomenon with a dramatic growth in the numbers of the ‘old old’ (and to a lesser extent of childhood disability) that has created the carers’ tipping point. In many western states, the increased numbers of disabled and frail elderly people has been accompanied by community living programmes. Whilst a debate exists as to whether the closure of large institutions can be attributed to human rights awareness or state ‘cost cutting’ the effect has been to increase the demand for ‘community care’. These two factors — demographic change and the welfare residualism that comes with neoliberalism — have resulted in a substantial increase in unpaid caring which in many developed nations, is nearing the limits of what families can provide.

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61 For example, The International Alliance of Carers Organizations and the European network organization EUROCARERS.

62 This paper uses neoliberal in its political sense: a system that adheres to civil and political rights and values, but believes that these cannot be used to limit the ‘free market’ — a natural organic entity that must be left, untrammelled, to flourish and so liberate individual entrepreneurial capacities and thereby create great wealth — see David Harvey, A Brief History of Neoliberalism (OUP, 2006).


64 As greater numbers of low birth-weight babies have survived — see for example, S Broach, L Clements and J Read, ‘Disabled children’ A legal handbook (Legal Action 2010).


66 Even if the ‘dependency ratio’ (the percentage of the population that is under 18 combined with the percentage that is over 65) is not itself changing dramatically — see N Folbre and J Nelson, ‘For Love or Money’ The Journal of Economic Perspectives, Vol. 14, No. 4 (Autumn, 2000), pp. 123-140 at 124.


However, at the same time in most OECD countries, there has been an even more remarkable increase in female employment rates.\textsuperscript{69} In the USA for example, women’s participation in the labour force has increased dramatically — for women age 55 and older the increase has been 50% in the past 15 years.\textsuperscript{70} During this period average household incomes have not increased\textsuperscript{71} - indeed they would have declined but for ‘women joining the workforce alongside their husbands’.\textsuperscript{72} For many women, much of the additional income from their employment is absorbed in paying for care costs — a process categorized as de-familialization — where a cost advantage (generally small) accrues by commodifying the care needs of both children and adult dependants.\textsuperscript{73}

Many of today’s carers, like the Vietnam veterans, are young and well educated, and aware that their adverse treatment derives from socio-legal environments constructed by and for people who do not have caring responsibilities: environments predicted on the ability to work and ‘inherently hostile’ to care-givers.\textsuperscript{74} Unlike the Vietnam veterans however, this group is predominantly female. It is an understanding of this question that produces the ‘creed’ – the necessary theoretical model that carers require in order to become a ‘rights movement’ – and which is discussed below.

\textbf{A narrative of oppression; a new historical account;}

There is considerable national and international evidence that carers in general experience adverse social, economic, health and political consequences as a result of their caring role.


\textsuperscript{70} MetLife, Caregiving Costs to Working Caregivers: MetLife Mature Market Institute, National Alliance for Caregiving, and Center for Long Term Care Research and Policy, New York Medical College (2011). This dramatic increase in female employment is mirrored in the UK

\textsuperscript{71} ibid


An impressive longitudinal well-being study of Australians found that female carers had the lowest collective wellbeing of any group it had considered and that Australian carers as a whole had an average rating that classified them as suffering ‘moderate depression’. Adverse impacts of this nature have been identified by a number of studies from Singapore to Greece, from Brazil to Italy and Norway. A 2011 study found that carers exhibit a higher prevalence of mental health problems across OECD countries than non-carers, with the rate increasing with the amount of caring. ‘High intensive’ caring in general increased the prevalence of mental health problems by 20%, but in Australia, the United States and Korea this became ‘70% or 80% higher’. UK evidence suggests that carers are a third more likely to be in poor health than non-carers and that over half of all carers have a caring related health condition for which almost 50% have sought medical treatment.

The severity of the adverse consequences experienced by carers is materially influenced by the nature of the state’s welfare support system. Thus carers in general are less likely to be in employment than non-carers — but this difference is less pronounced in modern welfare states (as for instance found in Nordic countries) than those with more residual systems. The prospects of employment bear directly on risks of poverty: working-age carers — particularly women — experience significantly higher rates of poverty.

76 EH Kua and SL Tan, ‘Stress of caregivers of dementia patients in the Singapore Chinese family’ (1997) 12 International Journal of Geriatric Psychiatry 466-469
77 Means and Smith cited above, p221.
In basic economic terms UK research found that at any one time a million carers have given up work or reduced their working hours to care\(^{88}\) and that as a consequence they were (in 2007) on average over £11,000 a year poorer\(^{89}\) that 40% of carers were in debt because of their caring roles (a figure that rose to 50% for parent carers)\(^{90}\). A 2011 study found that a third were unable to afford their utility bills and that three quarters had cut back on holidays, leisure activities, buying clothes and going out with friends and family\(^{91}\).

Research by the Australian Human Rights Commission has considered the long-term and gendered impact of the caring role. A 2009 study found that single elderly female households experienced the greatest risk of persistent poverty\(^{92}\) and a 2013 study\(^{93}\) found that the average superannuation payouts for women were little more than half of those for men. These differences were attributed to the struggle women experienced balancing paid work and caring responsibilities. Even discounting for the ‘accepted’ events in a woman’s lifecycle (pregnancy, childbirth and caring for children) the Commission considered that much of the difference stemmed from the ‘far less recognised’ caring relationships (predominantly) undertaken by women and which have the cumulative impact on lifetime earnings\(^{94}\). These findings are endorsed by USA research – that caregiving in early life significantly raised women’s poverty risks in later life\(^{95}\) and that older working caregivers had average pension shortfalls of $50,000 per person (in total amounting to a loss of nearly $3 trillion)\(^{96}\).

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\(^{88}\) Carers UK *The Cost of Caring* (2011).

\(^{89}\) *Out of Pocket, the financial impact of caring*, Carers UK, 2007


\(^{91}\) Carers UK *The Cost of Caring* (2011) and see also Carers UK, Carers in crisis (2008) and National Alliance for Caregiving and AARP, ‘Caregiving In the U.S.’ (NAC and AARP 2004) -0 where similar findings were noted in the USA research – p.13 and p62.


\(^{95}\) C Wakabayashi and K Donato ‘Does Caregiving Increase Poverty among Women in Later Life? Journal of Health and Social Behavior 2006;Vol 47: No. 3 258-274

\(^{96}\) *MetLife Balancing Caregiving with Work and the Costs Involved* (1999): a research study concerning people aged 50 or over, caring for their parents, undertaken by MetLife Mature Market Institute, National Alliance for Caregiving, and The National Center on Women and Aging.
Poverty is, as Alcock reminds us, ‘the unacceptable face of broader inequalities’ and in states with (or aspiring to) residual welfare systems, it is inextricably linked with employability. In such states the hostile nature of labour arrangements is a root cause of the adverse experiences of carers: operating as they do, in workplaces based on the notion of an ideal ‘autonomous’ worker – who it is assumed has “someone else” at home to raise his children. Work environments that ‘far from structurally accommodating or facilitating caretaking … operate according to premises that are incompatible with obligations for dependency’. Where:

Workers (at least some of them) must shoulder the burdens assigned to the family, while market institutions are relieved of such responsibility (and are even free to punish workers who have trouble combining market and domestic labor).

The politics of dependency

Dependency work (paid or unpaid) is gendered and it is this factor that lies at the heart of the injustice that carers experience. Caring is not, of course, an exclusively female activity – it is just that the status of caring has been engendered by the fact that it is women who provide the bulk of it. In the USA for example, it is estimated that there are over 25 million caregivers of which between 59% and 75% are women and that women on average spend 50% more time providing care than male caregivers. This is in line with the evidence from other OECD countries.
Whilst dependency is viewed as problematic in many political regimes — in the developed nations it is the neoliberal reification of individual independence, autonomy, and self-sufficiency that so disables and handicaps carers, as well as the people who depend upon them. These are values, which in Martha Fineman’s opinion have attained sacred and ‘transcendent’ status: but which are a myth: for ‘all of us were dependent as children, and many of us will be dependent as we age, become ill or suffer disabilities’. Dependency is hard wired into humanity: it may be a challenge but it is absurd to characterise it as unnatural — it is simply ‘inevitable’. The core creed for the carers’ movement is, therefore, the ‘politics of dependency’: just as we have created environments based on the needs of non-disabled people, so too have we created environments based on the mythology of independent people. Caring, like disability, is not in itself a handicap: it is the socio-legal context in which it is practiced that renders it so. The principal politically engineered handicap experienced by carers is that their care is uncompensated and — as Fineman and others have articulated so clearly — it is uncompensated because it is gendered.

Whilst the denigration of dependency and the marginalisation of dependency work is the aspect that most clearly explains the injustice that carers experience, a compounding role is played by the context in which it is generally practiced: the context of ‘privacy’.

A daunting literature exists that critically analyses the way that states have sought to create a socio-legal space — the space of the ‘private and the family’ — into which expansive ‘public’ notions of justice and equality should not intrude. The sphere of the ‘private’ (or what has been termed the ‘assumed family’) is an ideological construct that validates the severance of ‘individual dependency, pretending that it is not a public problem’; it is (Fineman once more) one that ‘masks the dependency of society … on the uncompensated and unrecognized dependency work assigned to caretakers’. This is, as Julia Twigg has described it, ‘dirty work’ and ‘hidden work’ — hidden:

because it deals with aspects of life that society, especially modern secular society with its ethic of material success and its emphasis on youth and glamour, does not want to think about: decay, dirt, death, decline, failure. Careworkers manage these aspects of life on behalf of the wider society.

107 Ibid p180.
108 see for example, Susan Moller Okin, ‘Justice, gender and the family’ (Basic Books 1989).
109 See for example Martha Albertson Fineman, ‘The Autonomy Myth: A Theory of Dependency’ (New York: The New Press, 2004) and Martha Albertson Fineman, ‘Cracking the Foundational Myths: Independence, Autonomy and Self- Sufficiency’, in Martha Albertson Fineman and Terence Dougherty (eds) Feminism Confronts Homo Economicus (Cornell University Press 2005) at 179 – 191; and see also Okin (citing Francis Olsen [CITATION – THIS IS AT PAGE 130 OF OKIN ] that ‘the very notion that the state has the option to intervene or not to intervene in the family is not only mythical but meaningless. In many ways the state is responsible for the background rules that affect people’s domestic behaviors’ Susan Moller Okin, Justice, Gender and the Family (Basic Books, 1989) p130.
110 Ibid (Cracking the Foundational Myths) at 179.
The creation of a different legal sphere from which many traditional legal principles are exiled, is essential to the maintenance of gendered systems. For neoliberalism, it is of particular importance since without the public/private delineation, key tenets would fall away. The dogma of ownership, for example – the right to own and sell the product of one’s labour – is self-evidently absurd when applied to a mother’s work in caring for her children. So too with commodification: if one commodifies caring – ie tots up the cost that carers should be paid for their caring work – then one ends up with very large sums indeed (‘unaffordable’ sums from a neoliberal perspective).

Rather than accept the severe limitations of such ideologies, a dustbin dimension is created – the ‘private’ space – into which all awkward facts a piled. States can then avoid accusations of injustice when failing to ensure that carers are properly compensated – and let individual carers bear this cost. Such an approach enables states to ‘ignore the crucial fact that much human labor, energy and skills is not devoted to the production of things that can then belong to their producers.

The radicalization of the carers’ movement is a consequence of the heavily gendered injustice at the heart of the current political settlement: a system that enriches those without impairments or caring responsibilities and consigns dependant people (children, elderly and disabled people) and their caregivers to poverty. A system that enables those without dependency to free-ride on the freely given care they received when dependant as children. For Fineman this injustice is currently ‘the most compelling’ problem facing our society: where ‘winners and losers become winners or losers in large part because of benefits and privileges or disadvantages and burdens conferred by family position and unequal distribution of social and economic goods.’

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112 In the UK a sum estimated as £119 billion pa L Buckner and S Yeandle Valuing Carers (Carers UK 2011) and in the USA using different criteria in 1997 it was estimated that the national economic value of informal caregiving amounted to $196 billion – see PS Arno, C Levine and MM Memmott, ‘The Economic Value of Informal Caregiving’ (1999) 18 Health Affairs 182-188 – but see also E Giovannetti and J Wolff, ‘Cross-Survey Differences in National Estimates of Numbers of Caregivers of Disabled Older Adults’ in The Milbank Quarterly Vol. 88, No. 3, September 2010: 310-349. Folbre and Nelson suggest that ‘valued solely on the basis of labor inputs’ it accounts between 40 per cent and 60 per cent of the total value of all U.S. output. As they observe, even this striking estimate contains a sizeable down-ward bias, since the market wages being imputed to women homemakers are lowered both by discrimination and by the time and effort put into nonmarket work’ - see Nancy Folbre and Julie A. Nelson, ‘For Love or Money’ The Journal of Economic Perspectives, Vol. 14, No. 4 (Autumn, 2000), pp. 123-140 at p 125-127

113 As Mitt Romney put it ‘inequality is the kind of thing that should be discussed quietly and privately’: cited by Joseph Stiglitz in The Price of Inequality (Allen Lane 2012) p 27.

114 Susan Moller Okin, Justice, Gender and the Family (Basic Books, 1989) p 129.

115 ‘Like other externalities, however, those created by care create an incentive to free ride, to let others pay the costs’ – see Nancy Folbre and Julie A. Nelson, ‘For Love or Money’ The Journal of Economic Perspectives, Vol. 14, No. 4 (Autumn, 2000), pp. 123-140 at p 137.

Caring and human rights

The language of human rights is the most obvious medium by which carers can express and challenge their collective marginalisation, and a number of authors have argued persuasively that there is indeed a human right to care.\(^{117}\) By conceptualising their situation in this way, carers can not only get closer to capturing the essence of their predicament,\(^{118}\) but they can also mobilise one of the few forces capable of tilting against the antagonistic political norms that manufacture their social exclusion. As West\(^{119}\) argues, it is only through the rights discourse that we can protect those facets of the human condition that we have come to understand as essential to our individual and collective ability to flourish and which ‘the political process is unlikely to confer on us’. In her opinion, rights are necessary:

> When for some reason, the sphere of life, service, freedom, activity, or identity that is protected by the right, and so necessary to flourishing, might nevertheless be systematically undervalued, underappreciated, or underprotected by standard political processes.\(^{120}\)

In the context of the struggle by disabled people, the articulation of a right to community (or ‘independent’) living is such an example. In many regions a good economic argument can be made in favour of institutionalisation and as a political issue, deinstitutionalisation is unlikely to be a priority for most voters. However, when articulated in terms of fundamental human rights\(^{121}\) the question is translated into an entirely new language – from one defined by the vocabulary of political and economic pragmatism to one of moral imperatives, urgency and repugnance. In precisely the same way, there is every reason to believe that until the adverse treatment of carers is understood as the proper subject of human rights, it will continue to be interpreted as a regrettable but economically inevitable fact of life.

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\(^{118}\) The ‘carers rights’ discourse has attracted a number of cautionary qualifications: Tronto for example suggests that it is within the idea of ‘an ethic of care’ that the analysis should take place U Tronto, ‘Beyond Gender Difference to a Theory of Care’ Signs: Journal of Women in Culture and Society 1987, vol. 12, no. 4 pp. 644-663 at p662) and Knijn & Kremer consider it better conceptualised as a dimension of inclusive citizenship (T Knijn and M Kremer, ‘Gender and the caring dimension of welfare states: towards inclusive citizenship’ in Social Politics (1997) Fall, 328–61 at 330). Important as these perspectives are, they do not undermine the idea that there is a ‘right to care’. Caring occupies a much larger and more profound space, than simply being a human right, but by so labelling it, it does not diminish this larger meaning: no more than referring to the right to religion in the language of rights can be said to restrict of compromise its value.


\(^{120}\) ibid

\(^{121}\) See for example, Article 19 UN Convention on the Rights of Persons with Disabilities and Article 26 Charter of Fundamental Rights of the European Union.
It has been suggested that a quasi-contractual / public law duty to secure compensatory support for carers is all that is required to address the impoverishment and hardships they experience: that the establishment of a right is superfluous. Of course the acceptance of a right to care, without a corresponding social support mechanism is of little value – but as West argues, persuasively, we need such a right ‘to protect caregivers against the pendulum swings of public support and neglect for their work’. Without such recognition, the carers’ needs would be ‘drowned in a tide of competing needs for scarce public resources’.

Arguably there are three (relatively) distinct dimensions to the human rights analysis and these will be explored in the succeeding section. The first concerns the proposition that there is such a thing as a substantive human right ‘to care’. The second concerns the extent of a state’s positive obligation to compensate carers for the adverse consequences of their caring roles. The third looks at the human rights of carers through the equality lens: that through this prism their adverse experiences can be seen as discriminatory.

Caring as a substantive human right

Civil and political human rights’ treaties protect various activities: expression; proselytising; marching / demonstrating; and so on. Although the essence of each activity has a platonic core, capturing this is generally problematic: political demonstrations are forms of expression; religious meetings necessitate association – and the extent to which these activities warrant protection can be graded in terms of their value: not all expression takes the form of the Gettysburg Address.

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122 See for example, Anne Alstott, ‘No Exit: What Parents Owe Their Children and What Society Owes Parents’ (OUP 2004).
124 Such ‘rights, if they exist, must be given content by legislatures through the normal mechanisms of democracy, not by courts through the extraordinary means of judicial review’ - Robin West A Right to Care in The Boston Review April / May 2004 at http://bostonreview.net/BR29.2/west.html accessed 10th June 2012
125 Other compartmentalisations have been advanced - for example, Stone argues for three facets of such a right, namely: (1) that families are permitted and helped to care for their members; (2) A right to care means, second, the right of paid caregivers to give humane, high-quality care without compromising their own well-being; and (3) a right to care must mean that people who need care can get it. See Deborah Stone, ‘Why We Need a Care Movement’ in The Nation, 13 March 2000, at 13 -14.
126 CHECK relevant ICCPR / etc Articles.
127 CHECK relevant ICCPR / etc Articles.
128 CHECK relevant ICCPR / etc Articles.
From a jurisprudential perspective, it is difficult to differentiate between the notion (or the ‘value’) of a right ‘to care’ and that of a right ‘to expression’ or ‘to belief’. All humanity arrives in this world utterly dependant and in need of care and for many, dependency is not a situation unique to their infancy. Caring has an elemental, non-commodifiable, altruistic quintessence,\(^{129}\) that is perhaps best characterised as a species of the fundamental human right ‘to give’:\(^{130}\) of a collective responsibility for dependency.\(^{131}\) It is the pre-eminent, indispensible and emblematic activity of a civil society – the absence of which is the acid test of inhumanity: of Brave New World and 1984.

To define ‘caring’- and hence the scope of the right – presents as great a challenge as defining ‘expression’ or delimiting the notion of ‘privacy’. Fundamentally, it involves providing care to meet the needs of a dependant person. The caring may be a physical, an emotional or a purely cerebral activity. It may involve intense intimate care: ‘dirty work’, ‘bodywork’ which may ‘involve inflicting embarrassing or painful procedures’… out of sight and in the back bedrooms’.\(^{132}\) Even if physical, it may be relatively impersonal – for example changing bedclothes or keeping a home clean for an elderly relative. It may be highly charged in terms of emotional support – of trying to keep a person from descending into depression; of ‘keeping their spirits up’; counselling and so on. It may consist of nothing more than ‘being there’ to ensure that the other person does not come to harm – of ‘keeping an eye’ on a young child or an elderly relative with dementia. Caring in this context is what the carer does – it is their physical or intellectual or emotional activity that makes it caring. The recipient may be grateful or ungrateful; oblivious, unconscious or simply indifferent: it is in this sense a classic gift relationship.

The person for whom the care is provided must have some element of need for that care. This may be due to the consequences of age (a young child or a frail elderly person) or that person’s impairment – be they mental or physical disabilities. Although the need may arise because of socially engineered barriers (physical, administrative, attitudinal for example), it is the need that is relevant – not its provenance, complex as this will sometimes be.

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\(^{129}\) Busby refers to the ‘intrinsically intimate nature of the exchange that takes place between a carer and a recipient of care’ that demonstrates ‘the inalienability of certain aspects. This central component of the relationship is crucial to the well-being of both parties and is, thus, non-commodifiable’ – Nicole Busby, ‘A Right to Care?’ (Oxford University Press 2011), p7; see also in this context Nancy Folbre and Julie A. Nelson, ‘For Love or Money’ The Journal of Economic Perspectives, Vol. 14, No. 4 (Autumn, 2000), pp. 123-140 at p129.


Whilst the scope of this paper is limited to ‘unpaid’ caring it is debatable whether the absence of remuneration (that the ‘work’ involved in delivering the care has not been commodified) is of pivotal relevance. Much has been written on this issue\textsuperscript{133} and the constraints of this paper enable it to avoid this contested and wide-ranging question.

That said, the mere fact that aspects of an activity are capable of being priced does not in itself render the process without value. Arguably commodification in the context of social care is better understood as an ethical or political discourse: tangential to the potential categorisation of the activity as a ‘human right’.\textsuperscript{134} Whether or not the process undermines and devalues is a mature debate – from blood donation to foster parenting: but the availability of blood engages a state’s obligations in relation to the ‘right to life’ and foster children’s relationships with their foster parents are categorised by courts as ‘family life.’\textsuperscript{135}

**The case for recognition**

Human rights treaties / constitutional provisions do not list every right of fundamental importance – for example the right to breathe\textsuperscript{136} or to feel the caring touch of the human hand. Instead, the relevant Articles are treated as living instruments within whose reach all rights essential to human flourishing are capable of being identified – be they (for example) the right to a livelihood,\textsuperscript{137} to development\textsuperscript{138} or to palliative care.\textsuperscript{139}


\textsuperscript{134} This issue was addressed by Munby J in *R (A and B) v East Sussex CC* [2003] EWHC 167 (Adm), [2003] 6 CCLR 194. at para 116 where he cited *Niemietz v Germany* (1992) 16 EHRR 97 at para [29] where the Strasbourg Court stated that there was no reason in principle why the ‘private life’ protected by article 8 “should be taken to exclude activities of a professional or business nature”.

\textsuperscript{135} See for example *G v E, a local authority & F* [2010] EWHC 621 (Fam) – a case in the High Court of England and Wales.

\textsuperscript{136} Indeed, Deborah Stone, in arguing for a ‘Right to Care’, states ‘Care is as essential as the air we breathe’ see Deborah Stone, ‘Why We Need a Care Movement’, *The Nation*, 13 March 2000, at 13 .


Whilst the formulation of some rights necessitates emphasis of provisions at the socio-economic end of the human rights spectrum, this is not the case in relation to the right to care. In this context, the Strasbourg Court’s approach to arguments concerning the existence of a ‘right to sleep’ and a ‘right to social interaction’ is informative for present purposes. Deliberate inference with a person’s sleep has been held to engage Article 3\ref{Ireland v UK 1978} (torture, inhuman and degrading treatment), whereas state sanctioned activities that interfered with individuals’ sleep have been held to engage Article 8\ref{Hatton v UK 2003} (private and family life). In similar vein, the deliberate inference with a person’s ability to interact with fellow human beings has been held to engage Article 3\ref{Keenan v UK 2001} whereas a state’s failure to take action to remove barriers that handicapped a disabled person’s ability to ‘participate in the life of the community’ has been held to have the potential to engage Article 8.\ref{Botta v. Italy 1998, Zehnalová & Zehnal v. Czech Republic 2002} This identification of such a latent right to community living within Article 8 (and in analogous terms, in the US by the Supreme Court\ref{Olmstead v LC 1999}) has of course been followed by its explicit recognition in the UN Convention on the Rights of Persons with Disabilities, Article 19.

The classification of a ‘right to care’ as a human right might be challenged on the ground that it is binary – involving as it always must, another. On analysis, however, conjoined rights are not unusual: the right to marry,\ref{Footnote to ICCPR / ECHR sources for this.} the right to associate\ref{Footnote to ICCPR / ECHR sources for this.} and indeed the right to family life\ref{Footnote to ICCPR / ECHR sources for this.} are not wholly egoistic or autonomous rights. Just as a right to care is contingent on there being a person ‘in need’ of care (or a ‘protected class’ in the language of rights) certain fundamental rights are contingent on another right being engaged – the so called parasitic rights, of which Article 14 European Convention on Human Rights (non-discrimination) is a classic exemplar.

Accordingly, a policy of treating family carers less favourably than non-family carers was held by the High Court of England and Wales to constitute differential treatment based on a family relationship – and (in the absence of justification by ‘counterbalancing factors of a compelling nature’) to violate Article 14 in combination with Article 8.\ref{R v Manchester City Council ex p L 2001}
Source / foundational human rights treaties

A right to care rests most obviously within the generic ‘right to private life’: Article 17 of the International Covenant on Civil and Political Rights, Article 8 of the European Convention on Human Rights (ECHR) and Article 11 of the American Convention on Human Rights. Whilst the Human Rights Committee and the Inter-American Commission on Human Rights have given only a limited steer as to their interpretation of the notion of privacy, this has been more than made up for by the Strasbourg Court which has described the notion of ‘private’ in the most expansive of terms: including a ‘person’s physical and psychological integrity’ for which respect is due in order to ‘ensure the development, without outside interference, of the personality of each individual in his relations with other human beings’. Thus sexual rights, environmental pollution, physical barriers to movement, access to files, the denial of citizenship, and information about one’s illness have all been held to come within its reach. In the context of the needs of disabled people, the Strasbourg Court has been particularly attracted to the notion of dignity – declaring that the ‘very essence of the Convention is respect for human dignity and human freedom’ (Pretty v UK 2002:65) and in Price v. UK (2001) Judge Greve considered that the measures necessary to ameliorate and compensate for the impairments faced by disabled people formed ‘part of the disabled person’s bodily integrity’ (ie their Article 8 rights).

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149 see for example UN Human Rights Committee General Comment No. 16 (1988) and S Joseph, J Schultz and M Castan, ‘The International Covenant on Civil and Political Rights’ (Oxford University Press 2004) chapter 16


151 Norris v Ireland 13 EHRR 186 (1988).

152 Hatton v UK (2003) (36022/97); Times July 10th


155 Whilst there is no right under the Convention to a particular nationality, an arbitrary denial of citizenship may in certain circumstances raise an issue under Article 8 – see for example, Kuric v. Slovenia (2010) Applic no. 26828/06 13th July 2010.

In *R (A and B) v East Sussex CC*\(^{57}\) the High Court of England and Wales was asked to give general guidance as to how local authorities should seek to resolve the relative interests of two disabled people (to be lifted safely and with dignity) and their paid carers (to avoid risks of injury from manual handling). In its analysis, the court had particular regard\(^{58}\) to the Article 8 jurisprudence of the European Court of Human Rights.\(^{59}\) Having identified the fundamental importance of disabled people being lifted safely and with dignity, Munby J (as he then was) observed that this needed to be put into context: the context that carers had corresponding rights. In his opinion such claims ‘are necessarily affected when the individual brings his own private life into contact or close connection with other protected interests’, adding (para 118):

I simply do not see how in this almost uniquely personal context persons in [the disabled persons] situation can seek to rely upon the rights afforded to them by article 8 without allowing that their carers have, at least in some respects, corresponding rights which have to be brought into the equation. If article 8 protects [the disabled persons] physical and psychological integrity – and it plainly does – then equally article 8(2) must … protect their carers’ physical and psychological integrity. And if article 8 protects [the disabled persons] dignity rights – and in my judgment it does – then equally article 8(2) must protect their carers’ dignity rights.

Having so determined, Munby J observed (para 120):

I recognise of course that the compassion of the carer is itself a vital aspect of our humanity and dignity and that at a very deep level of our instinctive feelings we value and need the caring touch of the human hand. … Even those who do not believe in any God know that a human being is more than a machine consisting of a few rather basic chemicals operated by electric currents controlled by some animalistic equivalent of a computer located in the skull – and that, no doubt, is why we have an instinctive and intuitive preference for the touch of the human hand rather than the assistance of a machine. As disabled persons or invalids our instinctive preference is to be fed by a nurse with a spoon rather than through a naso-gastric or gastrostomy tube.

At the very least the *East Sussex* judgment confirms that the caring role is the proper subject for human rights discourse. In so doing, it uses the elemental language of rights - that ‘the compassion of the carer is itself a vital aspect of our humanity and dignity’ – language indistinguishable from that deployed by the scholars considered above.

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\(^{58}\) Other provisions it considered to be of relevance included Article 3(1) of the Charter of Fundamental Rights of the European Union (the right to respect for… physical and mental integrity).

\(^{59}\) Notably *Botta v Italy* (1998) 26 EHRR 241.
Whilst this paper has focussed on the ‘right to private life’, it is not the only human rights provision that could be construed as protecting a substantive right to care. Nicole Busby\textsuperscript{160} for example has provided a convincing analysis on the relatively narrow issue\textsuperscript{161} of how a ‘right to care’ can be identified within European employment law, as a mechanism for reconciling the conflicts and adverse consequences experienced by those involved both paid work and unpaid care.

\textbf{Carers and the right to support the ‘doulia’ right}

Inherent within all human rights treaties\textsuperscript{162} is the obligation on states, not merely to refrain from interfering with the substance of the protected right (the ‘negative’ obligation) but also to take action to prevent the right being undermined. In the European context this positive obligation requires that states take effective measures to ‘secure respect for private life even in the sphere of the relations of individuals between themselves’\textsuperscript{163} and which may involve the implementation of domestic laws that provide the necessary protection.\textsuperscript{164} The positive and negative components, although subject to distinct jurisprudential criteria are seen as indivisible, and in this context Eva Kittay’s\textsuperscript{165} notion of reciprocity in caring is particularly apt:

\begin{quote}
Just as we have required care to survive and thrive, so we need to provide conditions that allow others – including those who do the work of caring – to receive the care they need to survive and thrive.\textsuperscript{166}
\end{quote}

\textsuperscript{160} Nicole Busby, ‘A Right to Care?’ (Oxford University Press 2011).

\textsuperscript{161} While acknowledging that such a right could also exist in EU law for those who ‘do not engage in paid employment’ – ibid p 11.


\textsuperscript{163} \textit{X & Y v. Netherlands} (1985) application no. 8978/80, para 23.


\textsuperscript{165} Eva Feder Kittay, Love’s Labor’ (Routledge 1999).

\textsuperscript{166} Ibid p107.
Kittay coined the word ‘doula’\textsuperscript{167} to explain the reciprocal nature of dependency in such cases – that without a positive obligation to support carers, those for whom they care ‘will continue to remain disenfranchised’ and their carers ‘will continue to share varying degrees of the dependents’ disenfranchisement’.\textsuperscript{168} For West\textsuperscript{169}, such a ‘right to provide care without risking impoverishment or dependency is comparable in importance and priority to the widely recognized core liberal rights of privacy, speech, property or contract’. For Fineman the relationship is best characterised as ‘derivative dependency’ - where one person ‘assumes responsibility for the care of an inevitably dependent person’. Her aim is to capture:

the simple point that those who care for others are themselves dependent on resources in order to undertake that care. Some of those needs are for monetary or material resources, whereas others are more related to institutional or structural arrangements.\textsuperscript{170}

Derivative dependency is, she argues:

culturally and socially assigned in an inequitable manner according to a script rooted in ideologies, particularly those of capitalism and patriarchy. These scripts function at an unconscious (and therefore unexamined) level, channelling our beliefs and feelings about what is considered natural and what are appropriate institutional arrangements.

Fineman argues that we share a ‘collective or societal debt’ for this fundamental caretaking role,\textsuperscript{171} and with it an obligation to challenge the prevalent socio-economic mores that are not only inimical to dependent people, but also to carers: that far from ‘structurally accommodating or facilitating caretaking’, societal institutions and workplaces ‘operate according to premises that are incompatible with obligations for dependency.’\textsuperscript{172}

\textsuperscript{167} From the ancient Greek word ‘doula’ which signified a female servant or slave.


\textsuperscript{169} Robin West A Right to Care in The Boston Review April / May 2004 at http://bostonreview.net/BR29.2/west.html accessed 10th June 2012

\textsuperscript{170} Ibid p.184.


\textsuperscript{172} Ibid p.183
What is being described here is a social model of exclusion – similar but even more subtle and ‘unconscious’ than that we have come to associate with the experiences of disabled people. In *Price v. UK* (2001) Judge Greve gave what has come to be considered a classic statement of the positive obligations owed to disabled people under civil and political human rights provisions – the duty to take action to ‘ameliorate and compensate for the disabilities faced’ to the extent that ‘compensatory measures come to form part of the disabled person’s bodily integrity’. In so finding, she noted:

The applicant’s disabilities are not hidden or easily overlooked. It requires no special qualification, *only a minimum of ordinary human empathy*, to appreciate her situation and to understand that to avoid unnecessary hardship … she has to be treated differently from other people because her situation is significantly different.

Although in contrast the handicaps and social exclusion experienced by carers are all too easily overlooked – this cannot in itself diminish a state’s obligation to take compensatory measures to address the injustice and marginalisation created by their derivative dependency. The fact that states with a neoliberal or similar gendered bias see this as the natural order cannot – in the court of fundamental human rights – be an adequate excuse. No more than it can excuse the exclusion of gay, illegitimate, or disabled people.

Kitty Malherbe has identified a number of human rights provisions that relate to the notion of a state’s positive obligations to mitigate the adverse consequences that arise from assuming a caring role. These include requirements in the UN Convention on the Rights of Persons with Disabilities on States to provide support for persons with disabilities ‘and their families’ for ‘disability-related expenses, including adequate training, counselling, financial assistance and respite care’ and that:

… persons with disabilities and their family members should receive the necessary protection and assistance to enable families to contribute towards the full and equal enjoyment of the rights of persons with disabilities.

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173 Application no. 33394/96; Times 13 August: 34 E.H.R.R. 1285.
175 Article 28(2)(c).
176 Preamble paragraph (x).
In addition Malherbe cites General Comment 5 of the Committee on Economic, Social and Cultural Rights\(^{77}\) which stresses the importance of ‘social security and income- maintenance schemes’ for persons with disabilities and then notes:

… the support provided should also cover individuals (who are overwhelmingly female) who undertake the care of a person with disabilities. Such persons, including members of the families of persons with disabilities, are often in urgent need of financial support because of their assistance role.

Also of relevance in this context, is the requirement in the UN Convention on the Rights of the Child\(^{78}\) that States should afford families ‘the necessary protection and assistance so that it can fully assume its responsibilities within the community’ for the care of their children.\(^{79}\) General Comment 9 of the Committee on the Rights of the Child\(^{80}\) (which concerns the support for disabled children) stresses the need for action to ensure that disabled children and their ‘parents and/or others caring for the child do receive the special care and assistance they are entitled to under the Convention’.

The reciprocal nature of carers’ and dependant people’s rights means that a failure to provide compensatory measures to enable the dependent person to live with dignity, may subject the carers to intolerable hardship, which itself can be articulated in terms of breaching their rights to respect for their private and family life and their right not to be subjected to degrading treatment. \(\text{R (Bernard) v London Borough of Enfield (2002)\(^{181}\)}\) concerned a claim by a disabled applicant and her carer that their human right had been breached by the failure of the local authority to take positive measures (by way of community care facilities) ‘to enable them to enjoy, so far as possible, a normal private and family life’. The claim succeeded because the council’s failure to act ‘condemned the claimants to living conditions which made it virtually impossible for them to have any meaningful private or family life for the purposes of Article 8’.\(^{182}\)


\(^{79}\) Preamble.


\(^{181}\) The High Court of England and Wales, [2002] EWHC 2282 (Admin); 5 CCLR 577; [2003] UKHRR 148, paras 32-33.

\(^{182}\) See also \(\text{R (Hughes) v Liverpool City Council [2005] EWHC 428 (Admin) 8 CCLR 243 paras 35 - 39}\) where a similar failure to provide support for a disabled person was held not to amount to a violation of that person’s ‘Article 8 rights solely because of the extraordinary efforts made by his carer (a burden the judge considered to be “intolerable”)’
Carers and Inequality

As noted at the outset of this paper, in the 1970’s the proposition that disabled people were the proper subjects of equality legislation was met with a degree of incredulity. However, within two decades their claim came to be seen as ‘self-evident’. Today a similar incredulity exists in relation to carers: being a ‘carer’, it is suggested, is not an innate characteristic and the handicaps they experience are those they assume when they choose to take on their caring roles.

Many carers do not articulate their experience in terms of choice: many speak of it in similar terms to the way disabled people describe their experience of impairment. The assertion of ‘choice’ does not of course vitiate the need for rational thought. Different societies offer different choices: being the parent of a disabled child or the child of a disabled parent is not a ‘choice’ and the options available to a person in this situation will be dictated in large measure by the welfare arrangements that the state chooses to offer. The situation has been described as ‘non-coerced yet not voluntarily chosen’ although ‘compulsory altruism’ is perhaps a better description.

The ‘non-innate’ argument is also suspect for a number of reasons, not least due to the existence in many states of legal obligations on carers to provide care and the moral coercion that exists in those others where no statutory liability remains. It is also undermined by protected status being accorded to ‘religious belief’: plausibly, it could be argued that ‘religious belief’ is no more immutable a characteristic than being a ‘carer’. Indeed, given the advances in medical technology (retina and cochlea implants, for example) the retention of disability has itself the potential to become a chosen characteristic.

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183 As Fineman observes ‘We ignore the fact that individual choice occurs within the constraints of social conditions. These constraints include ideology, history, and tradition which funnel decisions into prescribed channels, often operating in a practical and symbolic manner to limit options’ — see M Fineman, ‘Cracking the Foundational Myths: Independence, Autonomy and Self-Sufficiency’, in Martha Albertson Fineman and Terence Dougherty (eds) Feminism Confronts Homo Economicus (Cornell University Press 2005) at 185: or as George Elliot put it in Middlemarch, ‘there is no creature whose inward being is so strong that it is not greatly determined by what lies outside it’.


Indirect discrimination

It is arguable, therefore that the almost universal presence of legal, social and moral obligations on family carers has created a formalised public status of being a carer: one that transcends the private and the personal. Such a status may not be ‘innate’ but it is nevertheless very real. It is a role that falls disproportionately on women (expressly so in some states)\(^{186}\) and has undoubted negative health, financial, and well-being impacts.

In the language of existing human rights provisions such laws, domestic welfare arrangements and officially sanctioned social mores engage – indirectly, at the very least – several protected statuses: sex, disability, the rights of children for example, and in consequence demand of states, especial vigilance. Indeed such laws, policies and mores do more than ‘engage’ such statuses – they have a disproportionately adverse impact upon them, and constitute *prima facie* unlawful discrimination contrary to many international treaties: for example the International Covenant on Civil and Political Rights (ICCPR) (Articles 2 and 3); the African Charter on Human and Peoples’ Rights (Article 3); the American Convention on Human Rights (Article 24); and the European Convention on Human Rights (Article 14). Acknowledging this state of affairs, the Human Rights Committee referred to the ‘inequality in the enjoyment of rights by women [being] … deeply embedded in tradition, history and culture’ and stressed the need for States to ‘take all steps necessary … to put an end to discriminatory actions both in the public and the private sector which impair the equal enjoyment of rights.’\(^{187}\)

In similar vein, the UN Convention on the Elimination of All Forms of Discrimination against Women\(^{188}\) (CEDAW) requires states to promote measures which ‘enable parents to combine family obligations with work responsibilities and … participation in public life’ and which ‘eliminate discrimination against women in all matters relating to marriage and family relations’. The CEDAW Committee\(^{189}\) has noted that in ‘all societies women who have traditionally performed their roles in the private or domestic sphere have long had those activities treated as inferior’ and that ‘even where de jure equality exists, all societies assign different roles, which are regarded as inferior, to women.’\(^{190}\) At a regional level this has been echoed by the Council of Europe which has called for ‘the removal of barriers to positive parenting, whatever their origin’ and for employment policies that ‘allow a better reconciliation of family and working life.’\(^{191}\)

\(^{186}\) See footnotes 21 and 23 above [IRELAND / GHANA].

\(^{187}\) Human Rights Committee *General Comment No. 28: Equality of rights between men and women (article 3 International Covenant on Civil and Political Rights: 29/03/2000. CCPR/C/21/Rev.1/Add.10 paras 4 and 5*


\(^{189}\) Committee on the Elimination of Discrimination against Women, General Recommendation 16, Unpaid women workers in rural and urban family enterprises (Tenth session, 1991).

\(^{190}\) Committee on the Elimination of Discrimination against Women, General Recommendation 21, Equality in marriage and family relations (Thirteenth session, 1992)

\(^{191}\) Recommendation Rec (2006)19 Committee of Ministers on policy to support positive parenting, adopted on 13 December 2006.
In the absence of formal recognition – that it is unlawful to discriminate against a person simply by virtue of their caring status – such adverse treatment will fall to be characterised as indirect discrimination based on grounds of birth or sex. Indeed in relation to the preponderant role of women as carers, it has been argued, that there has been a ‘stunning silence’ about the effect of welfare policies on women, with the only debate being focussed on ‘how best to ensure such responsibilities were carried out’.

In response to claims by carers alleging unlawful indirect discrimination, states will be required to establish objective and reasonable justification for the difference in treatment. This in turn is likely to require evidence of the measures they have taken to ‘ameliorate and compensate carers for the handicaps they experience as a consequence of their caring role.’ In Strasbourg jurisprudential terms, the placing of status responsibilities of this kind, creates direct obligations on contracting states – as the court observed in *Marckx v Belgium*, a case concerning state policies which prejudiced the ‘illegitimate’ family:

> when the State determines in its domestic legal system the regime applicable to certain family ties … it must act in a manner calculated to allow those concerned to lead a normal family life!

**Associative discrimination**

Even if one accepts uncritically the argument that protected statuses should be reserved for those with immutable or ‘innate’ traits, there exists the challenge of the social impacts that are experienced by those in the out-group – ie people who associate with those who are protected.

A person may experience overt adverse treatment as a consequence of their ‘protected status’, without being subjected to explicit ‘direct discrimination’. In *Coleman v Attridge Law* (2008), a case before the European Court of Justice (ECJ) the Advocate General referred to ‘other, more subtle and less obvious ways’ – one of which was to target not the person with the protected characteristic (ie the black or disabled person) ‘but third persons who are closely associated with them and do not themselves belong to the group’. In the Advocate General’s opinion ‘a robust conception of equality entails that these subtler forms of discrimination should also be caught by anti-discrimination legislation’.

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193 Means, Richards & Smith, ibid p273.

194 Adopting the language used by Judge Greve in her concurring opinion in *Price v. UK* (2001) 34 EHRR 1285, albeit that the case related to a disabled person.


196 Ibid para 41.


198 Opinion of Advocate General Poiares Maduro delivered on 31 January 2008 Case C-303/06 S in Coleman v Attridge Law and Steve Law accessible at above web address – and para’s 12 – 14 in particular.
Coleman concerned the interpretation of an EU Directive\(^{199}\) which prohibited discrimination where a ‘person is treated less favourably than another’ on grounds of religion or belief, disability, age or sexual orientation. The applicant claimed she had been constructively dismissed for her employment because she had sought time off work to care for her disabled son: that her employer had treated her less favourably than employees with non-disabled children.

Her claim was problematical under the then UK anti-discrimination law\(^{200}\) since its prohibitions were limited to actions against ‘disabled people’ and it was the applicant’s son, not herself, who was disabled. The ECJ ruled however that she had been treated less favourably ‘because of disability’: that the Directive protected individuals from ‘associative’ discrimination of this type.\(^{201}\) The UK has since brought its legislation into line\(^{202}\) – and effectively carers are now protected from such adverse ‘associative’ treatment. A similar process has resulted in protection for carers in France\(^{203}\) and in Ireland.\(^{204}\) Protection of this nature is found (for example) in Peruvian antidiscrimination provisions,\(^{205}\) is under discussion in Australia\(^{206}\) and a duty to consider reasonable adjustments for working carers has already been enacted in New Zealand.\(^{207}\)

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\(^{200}\) The Disability Discrimination Act 1995.

\(^{201}\) Discrimination by association is not a new concept – particularly in relation to discrimination on the grounds of race: in the UK, for example, see Showboat Entertainment v. Owens [1984] 1 All ER 83.


\(^{204}\) Carers have formal protection (under the Family Status ground) in the Employment Equality Act 1998, s2(1) and the Equal Status Act 2000, s2(1).


Conclusions

This paper has sought to highlight the many parallels between the struggles waged by disabled people and by carers, in challenging their social exclusion. From the first domestic provisions addressing disability discrimination (most famously the USA’s Rehabilitation Act 1973) it took almost 30 years before concrete international legal provisions came on stream, such as the EU Equal Treatment Framework Directive 2000/78/EC and the UN Convention on the Rights of Persons with Disabilities. The period following the 1973 Act saw an exponential growth in states adopting specific disability discrimination legislation:208 action that formed the basis for effective universal recognition of the rights of disabled people.

We are now witnessing a similar ‘global’ trend in domestic legislation recognising the rights of carers. Carer specific provisions and ‘associative discrimination’ measures exist in almost every continent.

In Europe, for example, the EU has stressed the need for increased support for its 32 million209 ‘informal’ carers210 and for this to be put ‘at the top of’ each member state’s policy agendas.211 By 2009, Glendinning et al212 considered that in at least half the member states this had resulted in carer support being acknowledged as a political priority.

For neoliberal governments, in particular, the handicaps experienced by carers’ pose particular problems. Ideologically such governments espouse the ‘small state’ and are committed to a reducing the public provision of social welfare support. Unfortunately, in the developed Western nations this brand of economic liberalism has not (even before the financial crash of 2007) produced material benefits for the bulk of the population. As Stiglitz213 and others have observed, in such states a fall in middle class household incomes has only been averted by women re-joining the workforce. These changes have occurred at a time of dramatic increase in the numbers of dependent elderly people, and for whom institutionalisation is no longer considered appropriate. Carers — and they are preponderantly working women — are the elastic that has accommodated the contradictions in neoliberalism: a dogma that advocates work as the only route out of poverty but simultaneously holds to the belief that social care is primarily a family or charitable responsibility214. Carers are now stretched to breaking point, and these governments are aware of this.


212 Ibid

213 See footnote 72 above CHECK.

At the end of 2012, Cabinet papers from the first Margaret Thatcher administration were released, under the 30 year rule.\textsuperscript{215} Whilst these papers have attracted considerable publicity for different reasons\textsuperscript{216} what is striking is the Cabinet’s concern about sustaining ‘family caring’: its preoccupation with ‘the increase in the proportion of women’ in paid work; the ‘reduction in the ratio between the number of “typical carers” (women aged 45- 59) and the number of elderly people’\textsuperscript{217} and the ‘severe penalties’ that result from the ‘forces impelling women’ to take paid work.\textsuperscript{218} The policy direction of the government is summed up as ‘how to encourage families… to reassume responsibilities taken on by the state e.g. responsibility for the disabled… ’\textsuperscript{219}

As this paper has highlighted, the general response of governments since that time has been tokenistic: involving in large measure the enactment of opiate legislation – for example, of the relatively bland ‘carer recognition’ type. These are essentially rhetorical measures, heavy on process and exceedingly light on substance: responses that place little on no strain on the public purse. As Levitas has observed ‘recognising the value of unpaid work… means not recognizing its full economic value, since its cheapness is its main recommendation’\textsuperscript{220}

In his seminal paper concerning the impact of Americans with Disabilities Act 1990 Samuel Bagenstos (whilst celebrating the Act’s achievements) concluded that it had had ‘little, if any, positive effect on the overall employment of people with disabilities’ and little impact on eliminating ‘the deep structural barriers to employment that people with disabilities face’. In his opinion these problems could only be overcome by the government adopting ‘more direct and sustained interventions such as the public funding and provision of benefits’\textsuperscript{221}

For Fineman too, neoliberalism has no answers: the ‘approach to resolving this type of inequality is not found in simplistic and hypocritical prescriptions and ideological placebos of independence, autonomy and self- sufficiency’\textsuperscript{222}

\textsuperscript{215} The National Archives \textit{Prime Minister’s Office files} (PREM): Home Affairs. Family policy group; renewing values of society Catalogue ref: PREM 19/783 Date: 1982 May 26 - 1982 October 29

\textsuperscript{216} Not least for their revelation as to the chilling intent by that government to dismantle the welfare state – see Alan Travis \textit{Margaret Thatcher’s role in plan to dismantle welfare state revealed} The Guardian, Friday 28 December 2012 page 2.

\textsuperscript{217} The National Archives (Note 1? Above CHECK) at p89 - paper prepared by Secretary of State for Social Services (Norman Fowler) 7 September 1982.

\textsuperscript{218} The National Archives (Note 1?? above) at p148-149 - paper prepared by Secretary of State for Transport (David Howell) 27 August 1982.

\textsuperscript{219} The National Archives (Note 1?? above) at p12 - paper prepared by the Central Policy Review Staff FPG (82)2 November 1982.


\textsuperscript{222} Martha Albertson Fineman, ‘Cracking the Foundational Myths: Independence, Autonomy and Self-Sufficiency’, in Martha Albertson Fineman and Terence Dougherty (eds) \textit{Feminism Confronts Homo Economicus} (Cornell University Press 2005) at 189.
For carers and disabled people alike, the answer lies in the state providing decent support services for disabled people and by removing the barriers that handicap them. For this to happen a new political settlement is required: one that does not predicate everything on work – or work of the autonomous non-disabled model – but has at its heart, a progressive social welfare system.

Whilst the path that carers are treading towards the goal of a right to equal treatment is analogous to that taken by disabled people, it is not the same and indeed it has a different destination. The Disabled People’s movement seeks to create a society that is fully accessible and for which their different needs are accommodated and respected. Simplistically it is only if this struggle succeeds, that carers can have true equality: only when disabled people have full independence will carers have full equality.223

Simple as this assertion may be in theory – that if disabled people have a fully accessible environment and decent support services then carers will be able to have undisturbed lives – in practice it is not so. The reality is, of course, that the turbulent experiences of disabled people will always spill over and on to those close to them: it is the consequence of the human condition that we are affected by those closest and dearest to us. This impact (no matter how active, benign and universalist the state’s role) will always be capable of articulation in the language of disability and handicap, or alternatively in the language of experience and the loss of innocence. No state can compensate for such impacts since we will always have within ourselves an innate sense of our duty to care: a feeling that will inevitably open itself to exploitation by others – or indeed ourselves. A carer’s feelings of compassion, guilt and duty do not, however sanction adverse treatment, anymore than a woman’s maternal feelings justify treating her less favourably.

223 For those caring for children or frail older people, the ‘need is for a fairer society: one that does not just tolerate or (at best) ‘accommodate’ dependency, but one that regards it as central: as its raison d’être.