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System influences on work disability due to low back pain: an international evidence synthesis

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Abstract

Work disability due to low back pain is a significant global health concern. Current policy and practice aimed at tackling this problem is largely informed by the biopsychosocial model. Resultant interventions have demonstrated some small-scale success, but they have not created a widespread decrease in work disability. This may be explained by the under-representation of the less measurable aspects in the biopsychosocial evidence base; namely the influence of relevant systems. Thus, a ‘best-evidence’ synthesis was conducted to collate the evidence on how compensatory (worker’s compensation and disability benefits), healthcare and family systems (spouse/partner/close others) can act as obstacles to work participation for those with low back pain. Systematic searches of several scientific and grey literature sources were conducted, resulting in 1,762 records. Following a systematic exclusion process, 57 articles were selected and the evidence was assessed using a system adapted from previous large-scale policy reviews conducted in this field. Results indicated how specific features of relevant systems could act as obstacles to individual efforts/interventions aimed at tackling work disability due to LBP. These findings reinforce the need for a ‘whole-systems’ approach, with all key players onside and have implications for the revision of current biopsychosocial-informed policy and practice.
**Introduction**

Work disability due to low back pain (LBP) causes more global disability than any other condition [1]. In the United States (US) indirect costs of LBP are estimated to be more than US $50 billion per year, in the United Kingdom (UK) US $11 billion and in the Netherlands almost US $5 billion [2]. This burden is accounted for by approximately 10 percent of individuals with LBP, but the multi-factorial nature of work disability makes it very difficult to identify and thus prevent [3] [4, 5]. Research over the last two decades aimed at tackling this issue has shown that personal and occupational psychosocial variables play a more important role than spinal pathology or physical job demands; work disability due to LBP is now widely accepted to be a biopsychosocial phenomenon [4, 6-12].

The biopsychosocial model was developed in recognition that biological, psychological and social factors, and the interactions between them, can influence the course and outcome of any illness [13]. The approach was first introduced by Engel in 1977 [14] who stressed that the subjective experience of illness must be taken into account. The model proposes a dynamic systems approach, rather than one of linear causality or a factor-analytical approach as represented by the medical model. It also represents a shift of focus away from disease only, highlighting the importance of the illness trajectory in presenting opportunities to reduce/prevent disability. Although the International Classification of Functioning, Disability and Health (ICF) [15] acknowledges both the medical and social perspectives, it applies largely to individuals with impairments [16]. However, illness has many of the characteristics of a complex system, whereby dynamic interactions produce new properties and effects. Thus, the biopsychosocial model is particularly salient for understanding work disability due to LBP, which is now widely accepted to be the result of a trajectory of interactions between the individual and their social context [17, 18].

This understanding is reflected in the latest policy and clinical guidance for the management of LBP, which promotes early self-management and a continuation, or early resumption of, normal activities including work [19, 20]. However, the biopsychosocial evidence is dominated by research focusing on individual risk factors, despite the conception of the biopsychosocial model as a ‘whole-systems’ approach with all dimensions being equally important for work disability due to LBP [21]. Thus, resultant interventions are largely aimed at tackling unhelpful beliefs and behaviours, and whilst they have demonstrated some important successes, these are small-scale and have not created a widespread decrease in LBP disability [22]. Such individualised approaches to managing health conditions have been critiqued by not adequately taking into account the power that wider, systemic influences exert on individual will [23]. Yet research examining the non-modifiable
influences operating outside an individual’s perception and control, which cannot be adequately addressed by clinical and vocational rehabilitation interventions, is under-represented in the biopsychosocial evidence base [24] [25].

This issue reflects the perennial philosophical debate around the relative roles of structure and agency as they influence health [26]. The agency argument posits that health is influenced by the individual’s ability to act on decisions that arise from a unique self [27]. The structuralist view defines health as a product of context, and examines contextual factors that may impinge on individual behaviour, for example advice given by healthcare professionals [28]. To date, the biopsychosocial evidence in relation to work disability due to LBP appears to largely reflect the agency perspective, and has led to individualised interventions that are suboptimal.

In order to try and redress this imbalance in the biopsychosocial evidence base, this study provides a first attempt at collating and appraising the existing literature examining how relevant systems can act as obstacles to work participation for individuals with LBP. These systems are: compensatory (worker’s compensation and disability benefits – covering interactions with the workplace system), healthcare and family (spouse/partner/’significant others’) systems. Findings will have implications for the revision of current biopsychosocial policy and practice aimed at tackling work disability due to LBP.

Method
A ‘best-evidence’ synthesis was conducted. This was deemed the most appropriate method acknowledging that literature on the chosen subject is under-represented in the empirical evidence, and would be rather disparate and thus unfit for a systematic review [29]. A best-evidence synthesis gathers a range of academic (background and primary research, quantitative and qualitative) and grey literature available on the selected topic [30, 31], and draws conclusions about the balance of evidence based on its quality, quantity and consistency – it sets the results in context. This broad and inclusive method is particularly appropriate in order to capture the wide range of policies, practices, cultural tenets, and legislation in several different countries and settings. It also aligns to the methodology that has been successfully applied in relevant large-scale policy reviews which currently underpin the evidence base in this field [32-36].

The search strategy was compiled in an iterative process using both established MESH terms and keywords from the authors’ expert knowledge of the field. Although the primary focus of the review
was work participation and LBP, search terms related to other non-specific musculoskeletal conditions were also included as this evidence was deemed highly relevant to the study objectives. This is supported by the evidence which indicates that sub-grouping by non-specific musculoskeletal condition is unhelpful [37], especially when work participation is the outcome [36]. Conversely, it was decided that material relating to unemployed populations would be excluded because there is evidence that risk factors for work disability differ depending on employment status [4]. The included systems were identified as ‘compensatory’ (worker’s compensation and disability benefits), ‘healthcare’ and ‘family’ (spouse/partner/’significant others’)) due to their proposed importance as key stakeholders in the work participation process for those with LBP, but which had been under-researched [7, 38].

MESH terms and keywords, e.g. “Back Pain” or “Low Back Pain” and “Sick leave” or “Sickness absence” or “Absenteeism” or “Return to work”, and “Worker’s compensation” or “Sick benefits”/and “Healthcare professionals” or “Family Physicians” or “Health personnel” /and “Spouse” or “Social support” or “Significant others” were combined by Boolean operators (AND/OR/NOT), and various combinations within each category were attempted in each database so as not to miss any relevant sources. The searches were conducted in PubMed, CINAHL, EMBASE, PsychINFO, Scopus, Web of Science, Cochrane Library, ProQuest, Google Scholar and OpenGrey databases in December 2013. In order to maximise the chances of ensuring all relevant evidence was included across a range of disparate sources, no time limits were applied. However, only the evidence in English, Danish, Norwegian, and Swedish was selected due to the language comprehension of the authors. In addition to our systematic searches, we also included material recommended by experts within the field and a few additional evidence sources from weekly database alerts or authors coming across new publications were added concurrently until July 2014.

Evidence grading
A rigorous, yet pragmatic system was used to grade the available evidence (adapted from Waddell & Burton, 2006 [35] – see Table 1) whereby the grade refers to both the amount and quality of the evidence, indicating its ‘strength’. Due to the disparate nature of available evidence, it was not appropriate to apply a formal method of quality assessment to each article, and for the purposes of this study, quality was largely attributed to academic peer-reviewed published articles.

[Table 1 here]
Results

After removing duplicates, 1,762 records were retrieved. Initial title and abstract screens were conducted by one of the authors, the results of which were then discussed and agreed upon by all authors in an iterative process. Of these, further abstract/summary screenings were then undertaken by two of the authors independently, and 57 articles were finally selected for data extraction (see Figure 1).

Information from the included articles was summarised and examined [see attached supplementary material section for evidence tables], and findings are presented primarily in the form of evidence statements. These were developed, refined and agreed by the authors in an iterative process until consensus was reached. Each evidence statement was graded according to the nature and amount of relating evidence (see Table 1), and explicitly linked to the source(s) of this evidence. Evidence statements for each of the systems are presented as follows:

Compensatory systems

Evidence Statement 1:

* There is limited evidence to suggest that receiving worker’s compensation/disability benefit in itself, is an obstacle to work participation [39-44]

A critical review on this subject found that studies varied greatly in terms of methodology and design, with many having obvious flaws or shortcomings. [42]. Another review study indicated that some people view compensatory systems as a safety net that may reduce their perceived capacity, but this was regardless of health condition or not [44]. Thus, it was concluded that it was not possible to determine whether an association exists. For example, it was reported in one study that individuals had a significantly increased risk of prolonged claims if they had filed a worker’s compensation claim within 12 weeks of pain onset, therefore arguing for a negative effect of receiving worker’s compensation itself [39]. However, this effect was worse for those experiencing neck pain compared to LBP, and this difference was not explained. Similarly, another study reported a significant increase in return-to-work for non-compensated injured workers compared to compensated workers receiving functional restoration treatment, but this difference had diminished at 6 months [41]. In contrast, another study reported that receiving worker’s compensation did not affect time to return to work [43], and another found bi-directional effects depending on an individual’s health locus of control (the perceived level of control over own health condition). In
contrast to what one would expect, those workers reporting that the course of their pain was out of their control were more inclined to return to work compared to individuals who reported to be more in control [40].

**Evidence statement 2**

***There is robust evidence to suggest that specific, unhelpful characteristics of compensatory systems are obstacles to work participation [45-60]***

A wide range of studies contributed to this evidence statement, resulting in three common characteristics described as potential obstacles to work participation:

1) A strict and rigid compensatory system in terms of access to benefits and employment/modified work or worker’s compensation. The evidence here indicated that eligibility criteria should be less strict for long-term and/or partial disability benefits [53], but schemes providing widespread easy access to compensation were shown to discourage work participation, suggesting that unlimited compensation may have adverse effects [56, 58]. The presence of problematic rules and practices of the system, and of employers working against important initiatives like modified duties were emphasised [47, 50, 51, 60], and the rules and regulations of labour unions and compensators were specifically cited. It was reported that initiatives involving income insecurity during return-to-work are potential obstacles by reducing peoples’ incentives to engage in such initiatives [45, 46]. These were specifically described as comprising low benefit rates during re-education, a reduction in benefits when having some degree of work, and income uncertainty when trying to find a new job.

2) Specific elements of the compensatory system concerning regulation and conduct, such as right to case appeal, decision-making when medical uncertainty, or slow/dissatisfactory case management. There were some common features reported that seem to reward people for continued disability. For example, the access to an appeals process encourages an increase in labelling people as disabled because this achieves greater financial rewards. Furthermore, it was proposed that worker’s compensation/disability benefits relying on medical certainty complicate and prolong claims as individuals may be required to change course, be under constant medical treatment, and face defending their legitimacy through a cycle of claim denials and appeals [50, 51, 57]. Studies exploring the perspective of the claimant suggest that slow systems influence work disability directly or indirectly because they lead to a longer duration of work absence [5, 45, 49, 52, 59].

3) Regulations concerning the size of wage compensation rates following compensation claims. High wage compensation rates following injury claims were reported as an influence on work disability by increasing absence duration [48, 50, 55, 56]. Although one of the studies concluded that age and employer size were stronger predictive factors of prolonged absence than wage compensation rates.
[55], another did report a significant 1.6-increased risk of continued work absence for those receiving high compared to low wage compensation rates [48]. Wage compensation rates approaching or exceeding work wages were reported as a disincentive to work participation following injury claims [50], and that earnings-related compensation resulted in a slower exit of the compensatory system [56].

**Healthcare systems**

*Evidence Statement 3*

***There is robust evidence to suggest that a lack of work-focused healthcare (i.e. a failure by healthcare professionals to address work issues within the clinical encounter) is an obstacle to work participation [47, 61-81]***

Addressing work issues in the clinical encounter was defined as talking to the worker and the employer, posing questions and giving advice to the patient about work accommodation/date for return-to-work and prevention of re-injury, and referral to other relevant healthcare professionals. All except one of the studies relating to this evidence statement concluded that addressing work participation as a part of the clinical encounter would have a positive effect — the other study found that this effect disappeared when injury and workload characteristics were included in the analysis-model [61]. Reasons for not addressing work issues were most often reported as healthcare professionals not generally regarding engagement in work issues within their professional remit [66-70, 80]. Some of the reasons for this are proposed to include a lack of financial incentive [73, 74], a lack of time and standard procedures/role clarity, and increasing job demands [70, 73]. Some of the evidence indicated that healthcare professionals do not adhere to guidelines and give advice to (over)rest and take an unnecessarily long time off work [47, 79, 81]. It was suggested in one study that this was because healthcare professionals do not believe guidelines to be valid and therefore are unlikely to apply them [76]. In this vein, it was also reported that healthcare professionals’ distress about the complexity of LBP and their own ‘incorrect’ beliefs about LBP could be an indirect determinant for advising sickness absence and not engaging in discussions about work participation [66, 75, 78]. A seemingly important aspect was the difficulty healthcare practitioners reported with finding a balance between being a patient advocate and also gate-keeper of sickness absence certification. Many of the studies conducted in the countries where this kind of system is in place indicate that healthcare professionals allow a high degree of patient influence on their decision.
making about sickness certification in order to preserve the patient-practitioner relationship [65, 71, 72, 77].

**Evidence Statement 4**

***There is robust evidence to suggest that a lack of communication/cooperation between the healthcare system and other relevant stakeholders (e.g. employer, compensatory system) is an obstacle to work participation [44, 47, 63, 65, 80, 82-86]***

A lack of communication or loss of contact between relevant stakeholders [47, 80, 84, 85], poor communication skills, poorly communicated (and coordinated) work participation activities among stakeholders [82], and unidirectional communication between stakeholders [83] were specifically cited as influences on work participation for those with LBP. The evidence suggested that by improving communication between the healthcare system and other relevant stakeholders, workflow obstacles (time delays from incorrect or slow procedures) are seemingly reduced [80, 82]. In principle, additional issues such as conflicting demands from stakeholders [44] and lack of trust [80] can also be addressed. According to some of the evidence, a lack of common goals [47, 65], structural barriers, societal norms, healthcare professionals’ desire to maintain the professional status quo [83], and healthcare professionals being unaccustomed with involving others in their practice [44, 65] were proposed to act as barriers to communication. Where the effects of systematic cooperation between employer, occupational health service and social insurance office have been tested, a significantly immediate and sustainable effect on work participation (6 years) was found, but healthcare professionals highlighted difficulties due to “LBP patients’ inexhaustible needs for healthcare” [86].

Despite a general consensus among the studies that communication between the healthcare system and other relevant stakeholders was important, one study questioned the independent effects of this collaboration because they found that specific aspects of communication with healthcare professionals were more significant predictors for work disability [63]. Giving a patient a return-to-work date, and guidance on how to prevent recurrence and re-injury were each positively associated with work participation in this study. Furthermore, although healthcare professional contact with the workplace was found to be associated with work participation, this association became weaker upon adjusting for other variables (e.g. socio-demographic and job characteristics, pain duration and co-morbidity).
Evidence Statement 5
** There is adequate evidence to suggest that a lack of access to suitable and satisfactory health care is an obstacle to work participation [56, 87, 88]

Two of the studies relating to this evidence statement were based on workers’ perspectives, who cited geographic barriers to appropriate and timely care, and/or access issues related to workers’ “claimant status” (adding an extra administrative burden, reducing consultation time) [56], and satisfaction in terms of perceived quality of health care [87]. However, this latter study reported that an injured workers’ satisfaction with their employer’s treatment of their disability claim was more important compared to satisfaction with healthcare. Healthcare provided by the employer was reported to be important because it was suggested that not having access to health insurance, which provides easy access to healthcare, is an obstacle to work participation [88].

Family systems (significant others)
Evidence Statement 6
** There is adequate evidence to suggest that a lack of appropriate support from significant others is an obstacle to work participation [89-91]

Receiving support from significant others was considered to be an important factor for continued work participation for those with LBP, but the included evidence indicates that this is a complex interpersonal issue. For example, it was reported in one study that emotional support (availability of affectionately close and deep emotional relationships) increases self-appraisal and boosts self-esteem in individuals with LBP, but also encourages continued absence from work [89]. Another study reported that lack of support and distrust from significant others acted as an obstacle to return-to-work in persons absent from work due to LBP [90], and another found that solicitous behaviour from a partner increases disability levels among individuals with back pain [91]. Solicitousness was defined as getting the person in pain to rest, and taking over their jobs and duties.

Evidence Statement 7
* There is limited evidence to suggest that the beliefs of a significant others are an obstacle to work participation [6, 92-94]

An evidence-informed guidance document reported that negative beliefs, fears and expectations about work from a spouse/relative might serve as obstacles for work participation [6]. In addition, three small qualitative studies reported that significant others seem to share the negative illness
perceptions of their sick-absent family members, thereby indicating a likely detrimental influence. However, none of the studies investigated if these beliefs directly affected work participation [92-94].

**Discussion**

For almost two decades, a ‘whole-systems’ approach to tackling work disability due to LBP has been called for [60, 82]. This was reiterated more recently by The International Labour Organization (ILO) who strongly recommended that ‘enhanced social protection’ should be a key objective in the work disability research field [95]. Despite this, evidence examining the influence of wider influences has been lacking, i.e. the ‘social’ component of the biopsychosocial evidence base. The findings of this study make a valuable contribution to this evidence base, documenting how specific features of key (but under-researched) systems can act as obstacles to individual efforts/interventions aimed at tackling work disability due to LBP.

To illustrate this, we found limited and inconsistent evidence to suggest that receiving workers compensation/disability benefit in itself was an influence on work participation for those with LBP. Although there is some evidence to suggest that receiving worker’s compensation negatively influences recovery and vocational outcomes, largely due to the notion of ‘secondary gains’, the populations concerned are outside the remit of this review [96-98]. Moreover, a recent critical narrative review has dismissed such studies as biased because individuals with risk factors for work disability are over-represented among workers who file compensation claims [99]. Rather, it appears that a particularly important aspect is the timing of receipt of compensation/benefits in terms of making the transition less financially stressful. Regulations fostering employers and unions to indirectly discourage work participation are unhelpful - employers and unions need to convey that work participation is mostly the best outcome for the affected worker, providing that the job quality is good [35].

Lack of access to benefits during return-to-work, re-education or job transition were also identified as important influences - as was a claim being dependent on medical certainty or having access to an appeal process. Such regulations necessitate an open claim which prolongs and also reinforces disability; the evidence points to improved management of these processes. Again, this presents a challenge because non-modifiable factors, such as unemployment rates and the business cycle would need to be taken into account. Indeed, findings from an ecological study indicate that claim rate decrease in recessionary periods when unemployment increases [100]. Thus, our findings point
to the need for a ‘flexible’ compensation/benefits system with regulations that allow for adaptations related to the individual, the workplace and the business cycle.

The evidence also pointed to other salient issues such as lack of role clarity for relevant personnel, poor coordination of return-to-work activities, lack of knowledge and understanding of standard procedures, poor communication, and conflicting demands from other stakeholders. These factors were shown to result in slow or dissatisfactory case management, which in turn was suggested to prolong claims and delay return-to-work. In order to remove such barriers, the Organisation for Economic Co-operation and Development (OECD) has provided recommendations which result in financial stimuli for the relevant stakeholders (e.g. workplace, healthcare) to better collaborate. Such recommendations have been implemented in several European countries, and have been associated with significant reductions in sickness absence and disability claims for LBP [101]. More comparable studies need to be conducted, but the cultural and financial cross-country differences mean that this would be a significant undertaking. Such obstacles may explain the relatively lower number of studies focused on disability benefit/social security systems compared with worker’s compensation systems found in this review. Further research in this area is needed, and would provide a more nuanced understanding of how such systems affect work outcomes in populations with differing legislation, cultural norms, and socio-demographic statuses.

Moving onto the healthcare system, this review pointed to two particularly important process aspects: the practice behaviours of healthcare professionals and timely access to healthcare. Here, ‘practice behaviours’ refer to what healthcare professionals say and do to address work issues, communication and cooperation with the patient and relevant stakeholders (employer, compensation system, other healthcare professionals), and the patient-practitioner relationship. Robust evidence was found to suggest that failure to address work issues within the clinical encounter is an obstacle to work participation for those with LBP. The healthcare system is often the first point of contact for individuals with LBP, and it may also be the gatekeeper to other systems such as welfare and compensation. Healthcare professionals therefore, are in a prime position to administer the latest evidence-based guidance, which has recently called for healthcare to become more ‘work-focused’ [19] – that is, for healthcare professionals to take an interest in, and accept responsibility for, addressing obstacles to work participation in the clinical encounter. However, there is only sparse and conflicting evidence demonstrating the successful implementation of such guidance and its resultant effects on improving rates of LBP disability [102].
It is clear that further resources are needed to assist healthcare professionals to engage in work discussions with LBP patients, which can often be complex and challenging. For example, evidence shows that many individuals with LBP expect treatment to result in a significant reduction or complete removal of pain in order for it to be considered successful and for them to be able to work [103]. These unrealistic expectations and misconceptions are a difficult issue for healthcare professionals to manage, and often result in the use of practices that are not recommended and which can delay return-to-work, such as referral for imagery [104]. Such practices have also been proposed to result from misconceptions by healthcare professionals themselves about the work-health relationship [105]. However, the evidence in this study revealed a more nuanced and complex picture, and it would seem that in order to address non-adherence to guidelines, multidimensional initiatives and implementation strategies such as decision support systems; multilevel educational strategies; reminder systems; clinical practice audits; and regulatory change such as incentives to increased communication would be beneficial. Actions to ease the administrative burden on healthcare professionals to improve certain issues like time pressure, and access to proper assessment tools and communication platforms would also be helpful [106].

Findings also highlighted how the healthcare system acts a crucial stakeholder among various other important systems (e.g. the workplace and compensation/insurance systems). Thus, removing procedures that increase stakeholder conflict also appears to be of key importance. This is because healthcare practices that are unhelpful for the return-to-work process are often further reinforced by compensation and welfare systems requiring a medical diagnosis for a condition that most often does not have an equivalent pathology [107]. In the evidence reviewed, healthcare professionals described their cooperation with other stakeholders as unclear, scarce and often initiated late in the process. It was reported that healthcare professionals are generally not accustomed to initiating contact with other relevant stakeholders, and therefore it appears important to further promote a widespread understanding of the positive link between work and health amongst these stakeholders. Altering working procedures to include both relevant incentives and mandatory engagement and cooperation with other stakeholders, as well as improving the transparency and consistency of the return-to-work process is also required.

The other aspect revealed in this area of the evidence to be an obstacle to work participation was a lack of access to suitable and satisfactory healthcare, i.e. having access to work-focused healthcare at the right time. The retrieved studies indicated that lack of access was often due to simple geography or eligibility criteria related to ‘claimant status’, but a lack of health insurance through
the employer was also highlighted. Affordability has been identified as significant gap in health system coverage, and it is a problematic barrier in the context of most social welfare systems where equal access to health care is an overriding goal and principle [108]. One solution to these access barriers could be more provision of workplace ‘onsite’ healthcare, as supported in an international policy review which emphasized the value of investing in appropriate employee healthcare [109]. A way of increasing this is to legislate employers’ obligations to offer (suitable work-focused) healthcare to their workers the same way employers are already subject to legislation concerning workers’ safety. However, this would increase the financial pressure on smaller organizations, which may result in an adverse effect on worker well-being. This could be accommodated, however, by introducing tax exemptions related to workplace size or earnings/public budgets.

Another option would be if workplace healthcare access is considered a welfare task, and then it would be necessary to consider applying a more public health model providing case management and multidisciplinary support in the early stages of sickness absence, available to all. However, in order to create such a model, pilot studies testing the right construction of elements must be conducted. An example of this is the ‘Fit for Work Service’ introduced by the UK government [19]. The model proposes that when an employee has been absent from work for 4 weeks or more, the general practitioner (GP) refers the employee to an occupational health professional with the purpose of identifying individual obstacles for work participation and developing a plan containing recommendations and information on where to get appropriate help and advice. Employees, employers, and GPs also have access to advice from a phone line or website – this method of service provision is evidence-informed [33]. By introducing an additional tax exemption per year for each employee recommended to Fit for Work or an employer-arranged occupational health service, these are examples of how Government can influence a systemic and cultural shift in embedding work as a health outcome.

The final system proposed to be an important influence on work participation for those with LBP was the family system, or ‘significant others’. Although, it is widely acknowledged that interaction with spouses/family members is an important factor for recovery from pain, this was the most under-researched area compared with the other two systems in this review. A range of policy and empirical evidence has stressed the need to acknowledge the role of family members in order to improve health and work outcomes [110-115], but work participation is rarely the outcome in studies of this nature. Furthermore, data is rarely collected from family members themselves – it is either collected
from individuals with pain reporting their own perceptions of their family member’s beliefs and behaviours, or family member behaviours are observed and reported by a researcher or clinician.

Despite this, we found adequate evidence to suggest that a lack of appropriate support from the SOs of those with LBP influences their work participation. In this field, unhelpful support has largely been documented as solicitousness or punishment [116-118], and helpful support as empathic and positive responses [119-121]. However, the literature suggests that complexity can arise when this support or empathic response, viewed by couples as an indicator of relationship strength, translates into solicitousness which has a potentially detrimental effect [122, 123]. It has also been suggested that by providing help with everyday tasks, significant others are reducing the amount of activities for the person with pain, leaving more time for them to focus/ruminate on their condition. Furthermore, accepting this support may lead to the individual with pain to feel they are losing autonomy, thereby reducing their capacity to develop sustainable coping strategies. Conversely, providing (and describing) support may allow significant others to fulfil their ‘normal’ role as a caring family member in the face of an ‘abnormal’ situation [124].

This research illustrates the complexities which can arise due to the differing needs of those experiencing LBP and those close to them. The studies included in this review did not investigate such complexities, but our findings usefully point to areas warranting further investigation and highlight issues of practical concern in such research. Findings also indicate the potential of harnessing support from family members in order to improve work participation outcomes for those with LBP. This notion draws on the theory of ‘salutogenesis’ [125], which is in contrast to much of the research in this field which focuses on the negative or unhelpful factors involved. Indeed, it has been suggested that this ‘deficit’ approach may disproportionately lead to policies and practices which disempower those who are supposed to benefit from them by ignoring existing capabilities and capacities operating at an individual, group, community, population and institutional level that are protective or health promoting [126].

**Study limitations**

Compared with a systematic review, the method of ‘best-evidence’ synthesis inevitably involves a greater degree of judgement about article inclusion/exclusion, how the data were extracted, how evidence statements were developed and how the evidence was graded. While having to take much of the evidence at face-value, the significance of the various sources of data was weighted by virtue of its provenance. However, it is acknowledged that some relevant sources of international evidence
may have been missed due to the language comprehension of the authors. Such findings are more open to interpretation, particularly where studies were included due to relevance, and not only because they examined causal relationships with work participation. In order to minimise the risk of bias, the authors adhered to processes similarly applied in a systematic review, and described the process as explicitly as possible. However, it is acknowledged that the method is very inclusive and does not control bias as well as a full systematic methodology would allow, so the potential for bias is accepted.

Nevertheless, where pragmatic answers are needed to guide international policy and practice, we align with previous assertions that a broad and inclusive approach is needed [127]. This is because work disability research is ‘real-world’, social research and if a formal systematic review methodology was applied, then many relevant studies answering more pragmatic questions about non-clinical influences on work disability would not be selected [33, 128]. Acknowledging the importance of work participation for the health and wellbeing of so many individuals with LBP worldwide, we argue for a re-think of current hierarchies of study methodology that continue to exclude both the conduct and extraction of valuable, meaningful research demonstrating good work outcomes (or the potential of them).

This argument also applies to the use and acceptance of the biopsychosocial model in the field of work disability. The model has been criticised as failing to provide a more aetiological explanation as to which variables are the most salient, and whether these variables have equal weighting across all work-relevant health conditions and work participation statuses [129]. Indeed, the rationale informing this study was largely based on the acknowledgement that the biopsychosocial model has overemphasized individual, modifiable risk factors at the expense of other less amenable aspects. However, it is important to state that the biopsychosocial model is a process rather than a causal model, acknowledging that most illness is multi-dimensional. Thus, the salience of relevant factors may fluctuate at any given point, and this temporal aspect is a particularly prominent feature of work disability due to LBP [21]. Therefore, attempting to provide a fixed ‘weighting’ or ‘value’ to these factors would likely continue to obscure the ‘whole-systems’ nature of the problem.

This also lends itself to the view that work-relevant health conditions need to be understood and addressed using highly-specialised knowledge. Such an approach requires a high level of dependence on health and welfare services which will become unsustainable as the population (and workforce) ages [125]. It may be more helpful to appraise the different factors in terms of whether
they are amenable to individualised interventions delivered in the short-term, and those that require modifying at a policy level to effect culture change over a longer-term. This view would help shape our understanding that individual risk factors should be addressed as products of cultural and systemic structures that contribute directly or indirectly to work disability due to LBP [130]. Researching more of these upstream issues would hopefully inform the development of more accurate policy and practice, enabling all ‘key players’ to work together [131].

Conclusion
The findings of this study point to specific recommendations for the revision of biopsychosocial-informed policy and practice aimed at tackling work disability due to LBP:

- **integrating compensatory and health systems** to ensure individuals have access to what’s needed, when it’s needed, in a way which is personalised to their circumstances and needs;
- **embedding work as a health outcome** to stimulate all healthcare professionals to implement work-focused healthcare, and to promote the need for high quality occupational health provision, which will likely require changes in the educational curriculum;
- **ensuring a consolidation and standardisation of data collection around work participation** to facilitate future workforce planning and capacity requirements: these data will also facilitate a better understanding of what works for whom, when, and at what cost;
- **promoting the positive health benefits of work at a societal level** using a public health approach involving all ‘key players’.

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