ABSTRACT

Objectives This study aimed to identify priority policy issues and health system research questions associated with recovery outcomes for burns survivors in India.

Design Qualitative inquiry; data were collected through unstructured interviews and focus group discussions.

Setting Nine sites in urban and rural settings across India, through primary, secondary, and tertiary health facilities.

Participants Healthcare providers, key informants, burns survivors and/or their carers.

Results Participants acknowledged the challenges of burns care and recovery, and identified the need for prolonged rehabilitation. Challenges included poor communication between healthcare providers and survivors, limited rehabilitation services, difficulties with transportation to health facility and high cost associated with burns care. Burns survivors and healthcare providers identified the stigma attached with burns as the biggest challenge within the healthcare system, as well as in the community. Systems barriers (eg, limited infrastructure and human resources), lack of economic and social support, and poor understanding of recovery and rehabilitation were identified as major barriers to recovery.

Conclusions Though further research is needed for addressing gaps in data, strengthening of health systems can enable providers to address issues such as developing/providing, protocols, capacity building, effective coordination between key organisations and referral networks.

INTRODUCTION

Burns contribute significantly to the global burden of death and disability, resulting in at least 265 000 deaths annually and are one of the leading causes of disability-adjusted life-years in low-and-middle-income countries (LMICs). Burns consistently rank among the top causes of morbidity and mortality globally; however, have never received sufficient attention in global or national policy initiatives as they do not neatly fit under any single Millennium Development Goal and are not captured in the Sustainable Development Goals. WHO has, therefore, described burns as the ‘forgotten global public health crisis’.

In high-income countries, the incidence of burns has decreased and survival rates and recovery have improved, while in LMICs the incidence remains high and survival rates are poor. Burns are a striking example of the inequity of injury as the burden is borne primarily by poor people in poor countries where prevention programmes are uncommon and the quality of acute care is inconsistent at best.

Over half of all burns-related deaths in the world are in the WHO South East Asia region, where India has the largest burden with >1 000 000 people moderately or severely burnt every year, resulting in between 92 000 and 163 000 deaths. Further, it is estimated that 106 000 of these deaths occur in women, aged between 15 and 34 years. This age–sex distribution has been
consistent across multiple hospital-based studies, and the ratio of fire-related deaths of women to men is 3:1.7
Prevention efforts pose challenges particularly in the context of intent of the injury among women as self-imolation and domestic violence are common contributors to burn injury.8

Few patients with burns in LMICs receive appropriate first aid or immediate acute care, compounded by limited access to rehabilitation. In addition to physical disabilities, this often results in further complications such as anxiety, depression and post-traumatic stress disorder, often leading the patient to the conclusion that little or nothing can be done for recovery.9 As a result, burns survivors become emotionally overwhelmed and typically withdraw. Unfortunately, this lack of activity exacerbates secondary problems such as contractures, thereby heightening the survivor’s disability.10 11

The high burden of burns in India is known, and the challenges in prevention and care are recognised. However, there is poor understanding of the factors and interrelationships for planning effective measures for improving outcomes in burns survivors in India. The aim of this study was, therefore, to identify priority policy issues and health system research questions associated with recovery outcomes for burns survivors in India.

**Methods**

This study was informed by systems thinking, which views health in the broader dynamic social and economic context, recognising patterns and interrelationships with the aim of structuring those interrelationships in more effective, efficient ways.12 13

The research underpinned by systems thinking used participatory research for identifying priority areas for research, health system and policy aims to co-design interventions for the same. This study reports on the second phase of data collection in the participatory action research model. It informs on the networks, views health in the broader dynamic social and economic context, recognising patterns and interrelationships with the aim of structuring those interrelationships in more effective, efficient ways.12 13

Qualitative inquiry,14 consisting of interviews and focus group discussions (FGDs), was used to identify the needs of patients and carers within the burns healthcare context, as well as the experiences of healthcare providers and key informants, to develop acceptable approaches to address these needs in this setting. The rich diagram illustrates underlying mental models used during analysis. Rich diagrams (the precursor to Causal Loop Diagrams) are typically prototypes used to illustrate feedback and interactions among health system actors within a complex adaptive system.12 13 For this study, we present the initial identification and interpretation of the setting and interrelationship of factors that emerged in the context of burns care, influencing recovery outcomes in India.

**Setting**

India has a decentralised public healthcare system, run largely by its 29 states and 7 territories. The central government’s only constitutional mandate is to oversee medical education and collect infectious diseases statistics. In principle, coverage of health services in India is universal and available to all citizens under the tax-financed public system. Free primary care is provided by the public sector, supplemented by strategic purchase of secondary and tertiary care services from both the public and private sectors, which is the main financing approach.15

A National Programme for Prevention and Management of Burn Injuries was included in India’s 12th Five Year Plan in 2012.16 Increased resource contribution from the central and state governments was proposed. The programme commitment from policy makers holds promising opportunities for development of effective burns care system in India.16

This study was carried out across nine different sites in India, between June and October 2016. The aim was to include all primary and important secondary stakeholders at the meso and micro levels. A range of settings including urban and rural, health facilities ranging from tertiary to primary healthcare centres, private, charitable and public facilities and state settings using social development index as the primary indicator were included.

**Participants**

Participants in this study included healthcare providers, burns survivors, caregivers, neighbours/community, non-governmental organisations involved in rehabilitation, legal aid, workplaces and key personnel in health service implementation. Children were not included as participants; however, carers of child burn survivors were invited to participate. A purposive snowball sampling method was used whereby respondents were asked to suggest other potential stakeholders to be interviewed. This approach assisted with identifying and defining the boundaries of the network. Participant information is presented in tables 1 and 2, categorised as three main groups (patients and carers, healthcare providers, key informants).

**Data collection**

Prior consultation included in-depth interviews with healthcare providers only, and the key finding was the heterogeneity of healthcare services. The findings from the consultation helped identify key stakeholder groups and the breadth of healthcare settings. This paper presents the second round of data collection. We developed the interview and FGD guides based on a narrative synthesis of rehabilitation practices and recovery outcomes in burns survivors, and the initial stakeholder consultation. Before the interview or FGD, participants were provided standardised verbal and written information about the purpose of the research work, all data collection was face-to-face.
For this study, semistructured in-depth interviews and FGDs at burns care services in India were undertaken to explore the participants’ own framework of meaning; for example, what a good outcome means to a health professional and the injured person. Written consent from participants was sought prior to each interview or FGD. The interviewer used the interview or FGD guide (online supplementary appendix 1) covering a range of social, economic, cultural, psychological and environmental factors influencing burns care. Beliefs about

| Participant | Healthcare providers | Interview or focus group discussion | Type of facility | Years working in burns care/area |
|-------------|----------------------|------------------------------------|-----------------|---------------------------------|
| 1           | Social worker and psychologist | Interview | Charitable | 8 |
| 2           | Physiotherapist       | Interview | Charitable | 6 |
| 3           | General surgeon       | Interview | Charitable | 18 |
| 4           | Physiotherapist       | Interview | Public     | 5 |
| 5           | Nutritionist          | Interview | Charitable | 4 |
| 6           | Occupational therapist| Interview | Charitable | 5 |
| 7           | Nutritionist          | Interview | Private    | 3 |
| 8           | Plastic surgeon       | Interview | Charitable | 4 |
| 9           | Prosthetist           | Interview | Private    | 4 |
| 10          | Resident doctor       | Interview | Private    | 4 |
| 11          | Head nurse            | Interview | Public     | 12 |
| 12          | Nurse                 | Interview | Public     | 4 |
| 13          | General surgeon       | Interview | Public     | 14 |
| 14          | Emergency medical officer | Interview | Public     | 1 |
| 15          | General surgeon       | Interview | Private    | 11 |
| 16          | General surgeon       | Interview | Public     | 20 |

| Key informants | Interview | – | 7 |
|----------------|-----------|---|---|
| 17             | Paralegal  | – | 7 |
| 18             | State nodal officer | – | 2 |
| 19             | Non-governmental organisation representative | Interview | – | 7 |

| Patients | Gender | Interview or focus group discussion | Mechanism of injury |
|----------|--------|-------------------------------------|---------------------|
| 20*      | Patient| Focus group (nine participants of survivor group) | Mixed/unknown       |
| 21       | Patient| Interview                           | Flame burn          |
| 22       | Patient| Interview                           | Self-immolation     |
| 23       | Patient| Interview                           | Occupational—factory plastic |
| 24       | Patient| Interview                           | Occupational—gas service power station |
| 25       | Patient| Interview                           | Kerosene stove burn |
| 26       | Patient| Interview                           | Occupational—apprentice at electrical board |
| 27       | Patient| Interview                           | Occupational—contractor |

| Carers |
|--------|
| 28     | Carer  | Interview                           | Aunt of paediatric burn survivor |
| 29     | Carer  | Interview                           | Father of child who fell into fire furnace |

*Focus group discussion was all female participants and purposively male participants dominated the sample covering broader aspects other than gender and domestic burns.
practices, perceptions of their benefits and harms, effects of these practices and decision-making processes in health facilities, families and communities related to burns care were also explored.

Interviews lasted between 40 and 60 min in locations acceptable to the interviewees. Interviews were conducted in Hindi and Marathi, recorded, translated and transcribed in English. Handwritten field notes were made. To minimise data loss, interviews were jointly translated and transcribed by the interviewing researcher and the translator, and a member of the research team checked the translated transcripts with the Hindi and Marathi transcripts. Interviews ceased when saturation of meaning was considered to have occurred.17

The FGD was undertaken with nine burns survivors, caregivers and community workers. Participants were identified by the researchers in advance through patient support groups, healthcare providers and community workers. Participants were invited for discussion at a safe location in the community.

Data analysis

Transcripts were imported into NVivo qualitative data analysis software V. 11 (QSR International, 2015). Content analysis was applied to the transcripts whereby overarching themes were initially developed, informed by the understandings of a systems thinking perspective. Thereafter, deductive thematic coding of the data was undertaken. NVivo was used to arrange the text according to codes and manage the codes in the interpretive phase.

Through the qualitative data analysis, a rich diagram (figure 1) was developed using the software package Vensim PLE (Ventana Systems, 2015). For this study, the purpose of the diagram was to assist in the identification and interpretation of the context, components and outcomes that emerged in the context of burns care.

RESULTS

The interviews provided rich data, which described participant behaviour in relation to their service seeking, adherence to treatment and the broader sociocultural influencers throughout their burns care journey in the healthcare system.

Four themes were coded across all stakeholder groups, and resultant emerging threads were similar and interconnected. Quotes demonstrating these themes are

![Figure 1](https://example.com/figure1.png)

**Figure 1** System characteristics impacting recovery outcomes for burns in India.
DISCUSSION

The aim of this study was to identify priority policy issues and health system research questions associated with recovery outcomes for burns survivors in India. To contextualise our findings, we first present a discussion of the findings and how this was used to inform the development of the rich diagram (figure 1). We integrate this research into the wider context of burns, policy and health systems in India. Finally, this section will present considerations and research questions that emerged from this research.

Theme 1: shared tasks—care and rehabilitation

This theme articulates with the quotes presented in table 3, theme 1, and figure 1 section on patient flow in the healthcare system.

Participants from all three groups (patients and carers, healthcare providers, key informants) described the patient journey from the burns event through the healthcare system (figure 1—patient flow in the healthcare system). Limited resources were acknowledged but not seen as a major concern, with responsibilities shared between carers and health service providers. Time restraints and a shortage of health professionals willing to engage with burns care were raised. Participants, families and other carers were identified as playing an integral role as caregivers or attendees throughout the process of care. All groups raised the scope for their better inclusion in the care and rehabilitation process (figure 1—patient flow in the healthcare system).

Patients and carers expressed they experienced a gap in communication with healthcare providers, with many reporting not having been provided sufficient information about the care needed, and thus the level/type of care that they were able to provide in turn. There appeared to be a disconnect in healthcare provider perceptions around what information they provide with regards to rehabilitation education and practices, and this was particularly evident for information around burns dressings, dietary needs and posthospital physiotherapy, as well as resultant issues such as contractures.

Burns survivors and carers expressed frustration negotiating the complex, resource-intensive care system, with high out-of-pocket costs, access (transportation) issues and poor understanding of recovery. Patients and carers appear to trust healthcare providers; however, they also felt that they could not query healthcare providers around issues of care due to insufficient knowledge, indicating a potential cultural/education barrier. This suggests that there is scope for better processes around patient and carer education. One patient in particular suggested that people-centred care within the health system would be empowering (table 3, theme 1).

Theme 2: coordinated—multidisciplinary care

This theme articulates with table 3, theme 2, and figure 1 section on patient flow in the healthcare system, and overall strengthening health systems.

Stakeholder roles were varied, and the need for strong, cohesive multidisciplinary teams was well-recognised. Healthcare providers stressed that working in a multidisciplinary team is extremely important to the success of providing optimal care (table 3, theme 2). Healthcare providers often referred to burns care as a ‘not rewarding’ area to work in as recovery outcomes are often poor due to barriers, multiple levels of care and lack of multidisciplinary coordinated care.

The shortage of nursing staff and overall workforce (human resources) in burns care was evident. These healthcare providers expressed a great need for not only training but also retention of nurses (and other key hospital staff). As such, caregivers and attendants are regarded an integral part of the multidisciplinary team, which poses a question around better processes for their inclusion in the healthcare system. It is recognised that the quality of such assistance is variable, but also that better inclusion to this end would be valuable. This connects with the issues emerging in theme 1 around shared tasks in care and rehabilitation. Similar challenges are also experienced in rural and remote communities in high-income context.

The key informants stressed the importance of legal and financial assistance, particularly for patients with burns who have experienced gender-based violence (table 3, theme 2). The findings integrate with work from Morgan et al., who emphasise that health systems are not gender neutral; gender being a key social stratifier affecting health system needs, experiences and outcomes. Gender influences how people interact dynamically in complex, multifaceted and context-specific ways, reflecting varying interests, values and power, and therefore is at the core of health and health systems research.

Theme 3: community, access and health delivery systems

This theme articulates with table 3, theme 3, and figure 1 sections on the patient flow in the healthcare system, strengthening health systems, as well as the gaps elicited.

A major issue with hospital admissions process was patients moving between hospitals in the very early stages of their care (often taking days before proper care is received), as described by patients and their carers (table 3, theme 3). This intersects with known negative outcomes for recovery when adequate care is not received timeously. Difficulties with transportation to health facilities compounded this lack of access to healthcare in the acute phase of care.

Satellite centres or community-based rehabilitation for follow-up of discharged patients were regarded as essential as patients often do not return to hospital due
| Theme                                           | Quotes                                                                                                                                                                                                 | Participant                                                                                      |
|------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| 1. Shared tasks—care and rehabilitation       | ‘They are very important. Patient’s family has to give good support to the patient. If they would not maintain proper hygiene of patient, would not feed him nutritious foods, then his condition will not improve. We will do whatever is included in our list of duties but after that, only family members have to be there for him for 24 hours. While the patient is here, we are by his side 24 hours. Afterwards, someone has to be there with him.’  
‘Because you know we are not there 24 hours. The family members, one of the family members is there, so we have to explain things to him or her that why it is important how you have to feed the patient. This is very important.’  
‘Mostly, the health education is given by our nurses and doctors. And when the patient arrives, we explain things to the family members that time only.’  
‘If by chance, we give laundry in morning, if we cannot give it back in the evening. Earlier, we used to have only a single laundry system here. We ask them to use a clean sheet, iron it, and keep it with themselves. Give the patient a good diet. Never rush the patient. Do not ask him too many questions. For prevention of infection from mosquitoes and flies, use a mosquito net. Attend with clean hands. Sometimes, when we have too many patients, we give proper gloves to patients to apply ointment. They do it. It only happens when there is a huge number of patients and we tell our attendants to do the dressing of this patient and that patient too. We have to do it because there is a shortage of staff.’  
‘There should be one thing audio visual. Means audiovisual aids are the best and most need thing in our unit. We feel that in spite of telling them again and again they do not understand because they do not have that much education. We say “contractures” but they do not understand what that means. We use picture albums before discharge to reiterate the importance of rehabilitation.’  
‘We need to make tough decisions. We know which patients are salvageable, and who are not. Some are kept in general ward with just pain relief, some are sent back home. Burns unit has shortage of beds.’  
‘It is just that if they told us more then we could have made independent decisions.’ | Participant 12, healthcare provider, nurse  
Participant 7, healthcare provider, dietitian  
Participant 12, healthcare provider, nurse  
Participant 7, healthcare provider, dietitian  
Participant 2, healthcare provider, physiotherapist  
Participant 14, emergency medical officer  
Participant 25, patients and carers, male patient |
| 2. Coordinated - multidisciplinary team         | ‘We have not only read the importance of such a team in books.’  
 ‘So it’s a very nice multidisciplinary cohesive team where we all work together for improving the patient outcomes.’  
 ‘For any severe case of burn injury, an integrated team effort is required. Single doctor or single staff nurse cannot give full care to that person. Team and workforce is required. In our profession we learn about mental health psychological impact by experience, very late in the career. You cannot treat the body without treating the mind.’  
 ‘Burn cases are different. It is not like jaundice or pneumonia where the patient gets well fully and then, goes home. Like, I showed you pictures of burns cases. The patient could be experiencing (medical term), contractures, etc. So who will assess his mental status? For this, teamwork is required. And post-discharge, the patient requires physiotherapy. And occupational therapist will tell you about skills, etc. And, psychologists, counselors, and psychiatrists are very important. If these doctors are not on the team, and you send the patient home simply because his wounds have healed, then you will never know if the patient’s limbs are functioning well, if he has resumed his work, and if he is leading life as before. And, he will again try to commit suicide.’ | Participant 5, healthcare provider, nutritionist  
Participant 16, healthcare providers, general surgeon  
Participant 15, healthcare providers, general surgeon  
Participant 5, healthcare providers, nutritionist |
### Theme 3. Community, access and health delivery systems

| Quotes                                                                 | Participant                                                                 |
|------------------------------------------------------------------------|-----------------------------------------------------------------------------|
| 'I screamed and they took me to a hospital at [redacted]. It was a government hospital and it took too much time there. They kept asking me for my photo and papers. So much time was wasted there.' | Participant 23, patients and carers, male patient                            |
| 'They would ask you to deposit money first and then, they would begin the treatment.' | Participant 24, patients and carers, male patient                            |
| 'We had to move around 2–3 places, they kept saying take him there take him here.' | Participant 28, patients and carers, female carer                            |
| 'First we took him to the emergency. They cannot admit patients there. Then, we were sent to new PD (patient department) and two weeks later he was admitted.' | Participant 29, patients and carers, male carer                               |
| 'This can be done at the level of administrators in the hospital — my hospital and in hospitals, which are not related to this place like the various state governments, the people, who are sending patients to us. There, they could be doing a better triage, that send only those patients, who are really going to benefit. So, that is one. So the undue reference should not be there. Some patients could be managed only on advice, which can be done through tele-medicine, which we are now practicing. Like, many district hospitals in [redacted] will be sending pictures of their patients to us and then, we can discuss with them that don't send them here, manage them there. So, that's one way of utilizing your resources.' | Participant 8, healthcare providers, plastic surgeon                      |
| 'Follow up is the key issue because they don't want to come back to the same situation, but I think their lack of education and their lack of understanding leads to that kind of thing.' | Participant 6, healthcare providers, occupational                           |
| 'No, the [redacted] people had called me for a follow up but then, I did not have any money left. So I did not visit the hospital. Later on, in [redacted], I had to spend a huge amount. I talked to them about the expenses but they refused to do anything about it. Later, my father gave me money for my treatment. If I had money, I would have continued my treatment at the [redacted] hospital. They provided me with immediate care. But I had to arrange money.' | Participant 23, patients and carers, male patient, occupational             |
| 'I used to go to [redacted]. They used to call me and I visited them. If there were expenses involved then I used to take someone along with me. But I had to bear his expenses too. Like I had to provide him with breakfast, etc. If I promise to bear his expenses, only then he will come along with me. Moreover, I had to travel by special coach because with all this bulk, it was impossible to even think of travelling in general coach. People might push me or I might trip on something. So I used to travel by the special coach.' | Participant 24, patients and carers, male patient, occupational             |
| 'Yes, it would have been better, then there would be no need to travel always. They also asked me to visit on alternate days. But then, it would be too much for me. Travelling for 2 hours in heavy traffic would be so hectic. I don't have a car. And, it wouldn't be possible to travel by bus with all this.' | Participant 16, healthcare providers, general                               |
| 'Now the mobiles have come, we can do follow ups when there was no phones, there were no roads.' | Participant 24, patients and carers, male patient, occupational             |

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Continued
Table 3  Continued

| Theme | Quotes | Participant |
|-------|--------|-------------|
| 4. Systems—social, legal and policy challenges | 'How to handle the burns, what precautions should be taken when there is less burn, such awareness should be there. It is a developing country, in our industries, home, even in the hospitals there should be a separate department where patients should be told these precautions, how they should do the right things, and all this should be there … In my opinion, public awareness should be mandatory and the community should provide it in a timely manner.' | Participant 12, healthcare providers, nurse |
| | 'There is a stigma around burns. They do not want to be seen. We try but as you can see burns patients usually come to physiotherapy after all other patients. Spinal, geriatric, OPD (out-patient department) patients do not want to see them. It is sad, we try to educate everyone here but if there is stigma in health facilities you can see how hard it is for them to integrate socially. My friends tell me to work in sports injuries.' | Participant 2, healthcare providers, physiotherapist |
| | 'Burn awareness for prevention and minimization, prevention is one thing and minimization is one thing, should go hand in hand with the treatment also. Like, if they are establishing the units and all that and we are not giving the community the proper education about minimization or proper education about the prevention or how … So all these things should be told to the community or should be given to the community that these are the things you can do for prevention of this type of burns.' | Participant 18, key informants, nodal officer |
| | 'People should take care of safety at any place whether they are working for a company or otherwise. Initially, it feels difficult but all should adopt safety measures. One should be careful and companies should also practice it strictly.' | Participant 24, patients and carers, male patient |
| | 'See, we need to put these issues in front of government.' | Participant 6, healthcare providers, occupational |
| | 'Recovery is different, best in children. They are very well looked after too. Good thing is there is a shift, we do see male patients. Not all women as two decades ago, all dowry deaths. Now we have women 20 weeks pregnant, 3–4 pregnancy and you know why. Look around you' | Participant 13, healthcare providers, general surgeon |
| | 'The biggest barrier is that people are not interested. The burn in India is still treated as untouchable within the (unclear). In the sense that, although it does not happen in my unit, the burn patients are generally relegated to one corner of the hospital. They are considered unsuitable. But that mindset is changing at all levels. It never was in our hospital. Once you go to our hospital ward, you will see that our burnt patients are as neatly kept as comfortably kept as other normal patients. I think that untouchability part we are taking care of. They should be treated equally. But the mindset of people that burnt patients need to be treated with lot of care and compassion is not (unclear). Somehow, in this Indian subcontinent, we are not very sympathetic to burn patients.' | Participant 8, healthcare providers, plastic surgeon |
| | 'This is my personal perception, if this is the case, that is why the burns program is in a very initial stage and other programs are already taken ahead. This is the reason mainly because we have not considered because we have lots of other mortality and morbidity in India.' | Participant 18, key informants, nodal officer |
| | 'First thing is that legal rehabilitation is required. It should be this way that their cases will be given priority. Then preferably, their cases should be contested before female judges. Female judges are more sympathetic. And, they should be run fast, like fast track courts. Their judgments should come as fast as possible.' | Participant 17, key informants, paralegal |
to issues around understanding the need, as well as cost and convenience. Large gaps in follow-up after discharge therefore exist (table 3, theme 3). This intersects with the gap in self-care after discharge. Education around the importance of continued care is regarded as important. The most significant barrier to care is the cost (and inconvenience) of commuting for rehab and follow-up care (table 3, theme 3).

**Theme 4: systems—social, legal and policy challenges**

This theme articulates with table 3, theme 4, and the complete picture of the system depicted in figure 1.

The need for primary prevention of burns emerged as a key policy challenge. Healthcare providers further raised issues around government support and understanding of the issues, and education/knowledge translation as key policy challenges (table 3, theme 4). For the key informants, issues around national priorities for burns, and the need for prioritisation of burns cases in the legal system were other key policy challenges (table 3, theme 4). Social challenges such as gender, lower social economic status, poor education and stigma were identified. Health systems challenges of limited resources, out-of-pocket expenses also exist. These participants stressed that government support is needed to achieve outcomes around these aspects. Broader systems challenge of interpersonal violence, navigating the judicial systems, vocational training were the challenges raised by a range of stakeholders. This may be especially pertinent for survivors of burns as a result of gender-based violence where their support network is in jeopardy.

**Integrating this research with the broader context of burns, policy and health systems in India**

Based on the emergent themes above, and the interrelationship between key factors and stakeholders, figure 1 shows four key gaps that emerged out of the context of providing care and improving outcomes for burns survivors in India:

1. a structural gap, referring to discrepancies in terms of satisfaction of fundamental rights—access to healthcare;
2. a participation gap, referring to inhibitors to genuine participation in society—right and access to work, healthcare;
3. a feeling/aptitude gap, referring to psychological and social traumas, shame and mistrust;
4. a knowledge gap, referring to limited knowledge on, for example, communication—health/administrative regulations.

In the context of strengthening health systems, figure 1 therefore elicits opportunities for improvement around supply chain system design (efficiency, agility, resilience, responsiveness, execution), last mile availability (infrastructure and processes), resource mobilisation and supply chain operations financing (sufficient funding and distribution), governance and accountability (formal and informal incentives in the public health supply chain) and sustainable human capacity (developing, attracting and sustaining staff).17

It is evident from this study that care and rehabilitation services for burns survivors in India do not exist in isolation; rather, they play an integral role within the healthcare continuum and therefore must be considered within the context of the overarching health system and cultural context. People’s behaviour and experiences in this healthcare system can be explained by social, economic, cultural, psychological and environmental factors, which impacts on service-seeking behaviour, adherence to treatment and the broader sociocultural influencers.

Functioning of an effective health system can strengthen the capabilities of individuals and social groups, for example, by including people in identifying health priorities and planning services from individual care to community health interventions. People-centred health systems, in this way, reflect a progressive shift that has moved thinking beyond building-blocks models of health systems towards ones that centralise a human and relational nature. These approaches can also generate preferential gains for socially disadvantaged groups, either by impacting on the structural factors that disadvantage them (women’s autonomy) or by strengthening their ability to claim health resources or implement health action.

However, in practice, severe bottlenecks in accessing government healthcare services compel households to seek private care, often resulting in high out-of-pocket payments. Significant inequalities with respect to healthcare access and health outcomes exist between states, rural and urban areas, socioeconomic groups, castes and genders. Past decades have seen several initiatives targeted at improving primary healthcare systems; however, the focus has been on infectious conditions and maternal and child health.

Systems thinking approaches12 13 provide guidance on where to collect more data, or to raise new questions, hypotheses, obtain buy-in from key stakeholders and develop interventions that are pragmatic. A frustration for health planners and researchers has been the aspiration that interventions shown to be effective in a research setting cannot be simply replicated at large scale or to reach populations that are most vulnerable. Therefore, this research was conceptualised with the aim of eliciting a range of priority policy considerations and future research questions from a local, setting-specific and contextualised perspective. These are presented in box 1.

**LIMITATIONS**

Qualitative inquiry is time and context dependent. Therefore, this study is bound by the experiences and beliefs of these participants alone, but does resonate with what is known about the larger context. Furthermore, it was not possible to elicit relationships between the themes and the specific outcomes, although this is clearly an important area for further research. As the aim of this study was to provide considerations for future research, the method

Jagnoor J, et al. BMJ Open 2018;8:e020045. doi:10.1136/bmjopen-2017-020045
Documenting the burden and recovery outcomes among burn survivors

- Scoping burns registry to document burden and improve pathways for care for better recovery outcomes.
- How can health-related quality of life for burns survivors in India (low-and-middle-income countries) best be understood and meaningfully captured? How can the outcomes for burns survivors be improved as they move through the health system?
- How can we evaluate the financial impact/financial disability among survivors of burn injury?

Quality improvement in care, equity and access in burns care

- Need for development of best practice guidelines which include rehabilitation prescription facilitating continuum of care for improved recovery outcomes.
- How can communication about patient care throughout their flow through the healthcare system and into community- and/or self-care be enhanced? What are the best options for improved knowledge translation?
- Focus on implementation research using real-world scenarios.
- Is there scope for mobile or digital health technologies to improve patient care throughout the healthcare system? What technologies exist? How can these be harnessed?
- Can (or should) task shifting to caregivers and family carers be formally included in a person-centred healthcare system? If so, how is this best achieved? What can be the pathways, family-centred care or by enabling community health worker?
- Adopting policy changes on health insurance and universal health coverage to address these challenges.

Systems approach to reducing burn injury-related burden

- Develop a systems framework for burn care and prevention using the WHO system building blocks incorporating systems organisation, networks, dynamics and knowledge.
- What is the current state of legal and financial support available for burns survivors? How is this communicated and administered? What steps can be taken to improve this process, particularly in the context of culturally sensitive issues around burns survivors such as gender-based violence?

CONCLUSION

The ‘invisible face of burns’ is a critical public health problem in India. A goal of this study was to feed into the future development of pathways of care for burns survivors using a systems thinking approach in a realist context. This research will enable development of acceptable models of healthcare that can then be developed and piloted in the next stage of the research process. Better understanding how and where to potentially influence optimal long-term recovery outcomes, including a return to activities of daily life and quality of life, through considerations of opportunities and gaps is a key contribution of this research.

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