Delayed Presentation in Leprosy

How to identify the causes

A guide to fieldwork
Acknowledgement

This booklet is part of instead of this research is part of the ILEP Nerve Function Impairment Research programme (INFIR), addressing issues of prevention of disability in leprosy.

We are grateful to the Raoul Follereau Association of Luxembourg and the Luxembourg Government for their funding of this research.

We also acknowledge the support and co-ordination role of the INFIR Steering Committee and the staff of the Leprosy Relief Association (LEPRA).
Correspondence:

Peter Nicholls, Department of Public Health,
University of Aberdeen, Polwarth Building, Foresterhill,
Aberdeen, AB25 2ZD, Scotland, UK

The procedures described here are drawn from published texts.
Every effort has been made to ensure the information presented is accurate.
The authors can accept no responsibility for problems that may arise in their use or for the costs incurred.
Illustrations by LJ Smith
Identifying the causes of delayed presentation in leprosy

A guide to field work

Purpose

This guide to fieldwork identifies basic fieldwork methods to investigate the reasons why people affected by leprosy delay in presentation and start of treatment. We describe the actions needed to prepare and supervise fieldwork, to analyse data and to prepare recommendations. In each chapter we provide some background information and identify recommended actions. Keeping in mind the needs of those whose first language is not English we have tried to keep the wording as simple as possible. At the end of the booklet are a series of Annexes providing more detail.
## Contents

<table>
<thead>
<tr>
<th>Chapter One</th>
<th>Introduction and Outline</th>
<th>Page 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chapter Two</td>
<td>The help-seeking process that adds to delay</td>
<td>Page 3</td>
</tr>
<tr>
<td>Chapter Three</td>
<td>Preparing for fieldwork</td>
<td>Page 7</td>
</tr>
<tr>
<td>Chapter Four</td>
<td>Learning from patient records</td>
<td>Page 11</td>
</tr>
<tr>
<td>Chapter Five</td>
<td>Learning from patients</td>
<td>Page 14</td>
</tr>
<tr>
<td>Chapter Six</td>
<td>Learning from the community</td>
<td>Page 21</td>
</tr>
<tr>
<td>Chapter Seven</td>
<td>Learning from staff</td>
<td>Page 27</td>
</tr>
<tr>
<td>Chapter Eight</td>
<td>Analysing your data</td>
<td>Page 33</td>
</tr>
<tr>
<td>Chapter Nine</td>
<td>Applying your results</td>
<td>Page 36</td>
</tr>
<tr>
<td>Annexes</td>
<td>Stigma, Fieldwork Skills, Action Plan, Methods, Evaluation</td>
<td>Page 39</td>
</tr>
</tbody>
</table>
Q. What is delay in presentation?

- For each patient, delay is the time from becoming aware of the first symptom through to start of treatment.

Q. How do I know if there is a problem?

- People who delay are more likely to develop impairments – nerve damage with sensory or motor loss or visible impairments such as ulcers. Do you have people like this in your project?

Q. Why should I get involved?

- Delay prolongs the time when a person may pass on the disease to others.
- Delay may mean more drugs and a longer course of treatment.
- Delay increases the risk of permanent disability.
Q. What would be involved?

To understand why people delay you need to understand how people assess their symptoms and the actions they take to seek help. These all contribute to delay. This will involve:

- Taking time to listen to the experience of individual patients.
- Learning from significant people within the community, for example, teachers or religious or political leaders.
- Talking to doctors involved in private or general health services.
- Hearing what your staff have to say.
- Preparing a summary of the primary factors leading to delay.
- Identifying the actions that may encourage early presentation.

Q. I already have enough work. Is this really important?

- Achieving early presentation is essential to controlling leprosy.
- The local reasons for delay are unique to your situation. No one else can do it for you.
- Your patients already know why they delayed. You can learn from them.
- Your staff and community members can contribute.

We are not suggesting a formal research project. The procedures we recommend rely on a few basic skills to produce the information you need to solve a problem. To get a clear picture, you should first read through this booklet.
Chapter 2: The help-seeking process that adds to delay

Figure 1 describes how the actions people take add to delay. They assess their symptoms, decide on an action and assess the result. While they rely on ineffective treatments or fail to suspect leprosy a series of such actions adds to delay. Finally, something happens to break the cycle. The symptoms are recognised. They attend the clinic, are diagnosed and start treatment.

Figure 1. Help-seeking actions that add to delay
Q. If this is why people delay, what is the problem?

The problem is in the great variety of ways in which people decide to seek help:

Decisions are affected by awareness of leprosy:
- People may not recognise the early signs.
- They may be unaware of the cure.

Decisions are affected by local culture:
- People may not be able to go for help without the approval of the husband, father or head of family.
- People may not recognise the early signs of leprosy as an important disease.
- People may prefer to use traditional medicine and cures.
- People may follow advice from owners of medicine shops or community or religious leaders.
- People may believe leprosy is inherited or caused by a curse or judgement, so no medicine can be effective.
Decisions may be affected by questions of cost.
- People cannot afford the loss of earnings during clinic visits.
- People may think the cost involved in going to the clinic and getting treatment is too high.
- People may think the consultation fee with a doctor is too high.

Decisions may be affected by fear of the social consequences (Annex 1).
- People may fear the consequences for their marriage or family or work.
- People may fear the consequences for their position in the community.

Decisions may be affected by experiences within the health services
- People may go to doctors expecting an instant cure. When this doesn’t happen they try a different doctor. They don’t give the doctor time to decide a diagnosis.
- People may reject the diagnosis and treatment they receive and look for a more acceptable diagnosis from another doctor.
- People may ignore the doctor’s referral to a specialist.
- The doctor may misdiagnose.

There are hundreds of reasons why people delay!
The reasons people present and start treatment are also of interest:

- They present because they recognise the possibility of leprosy.
- They present because their symptoms can no longer be hidden.
- They finally decide to follow the advice of family members, friends or doctors.
- They are found through survey.

In any given community some reasons for delay are more common or more important than others:

- It may be that the problem occurs amongst older people who maintain traditional beliefs.
- The problem may be among young women who don’t want to spoil their marriage prospects.
- The problem may apply in limited geographical areas.
- The problem may be amongst particular occupational groups.

Some basic fieldwork will help you locate and understand your local problem. The information you collect will guide the decisions you take to address the situation.
Chapter 3 - Preparing for fieldwork

Methodology

• Rely on interviews, group work and observation. These methods are best for exploring new areas and uncovering new facts and practices. Chapters 5, 6 and 7 each describe one method. See also Annex 4.

• Your findings must not be biased by one point of view. Plan to work with patients, and with at least one other source of information – your staff, health service practitioners or individuals or groups representative of the community. We also recommend you learn from the data stored in your patient records. You can be confident of your findings when two or more sources or methods give matching information.

• If you follow the suggestions in each chapter you should be able to complete the whole project in 8 to 10 weeks. In the outline Action Plan (Annex 3), we propose three phases of work, two weeks of preparation and training, six weeks for fieldwork and one to two weeks for analysis and report preparation.

• Don’t wait until the end of fieldwork to look at the data - instead review the data as it arrives. You may find your fieldworkers need more training in the chosen methods. It may be you need to set a new focus and ask some extra questions in some of the fieldwork. You need to make these changes as soon as possible.
Make the most of the people and resources available to you:

People:
- Recognise patients and staff as important sources of information.
- Identify key individuals or groups within the community as additional sources.

Data:
- Make use of data already available to you – specifically, your patient records.

Skills:
- Make full use of existing skills amongst your staff, in particular, their interviewing skills.
- Involve local organisations that can offer skills or other forms of assistance.
- Remember that practitioners in traditional medicines have the best insights into local health practices.

Funds:
- To save on travel costs, plan fieldwork in and around regular outpatient clinics.

Annex 2 contains more information on fieldwork skills.
Keep the work to a manageable minimum:

- Choose to focus on a limited geographical area.
- Choose to focus on specific socio-economic groups
- Choose to focus on a specific demographic group.

You can always return to other groups later.
Q. Am I really going to learn anything new?

- Keep focused, but also keep an open mind.
- Don’t devalue the experiences of your patients. Their stories provide the information you need. Try to understand their thinking and decision-making.
- Respect the point of view of your staff, especially those with long service. They have had contact with many patients, in the clinics and in the homes. They have learned something about delay.

Q. Will anyone take any notice?

- Your manager must be convinced there is a problem and ready to respond to recommendations you make.
- Be sure your manager approves of the fieldwork.
- Involve your manager in progress reviews, problem solving and field visits.
- Maintain his/her involvement through reports and in identifying recommendations.
Chapter 4 – Learning from patient records

Who are the people that delay?

The first action is to identify and describe the people who delay in presentation or first present with visible impairments. To do this we will collect information from the patient clinic records:

- Work through the most recent registrations, people registered in the last 3 to 9 months. Earlier registrations may not be relevant to the current situation.
- If you are focussing on a limited geographical area or a small patient group, choose your records accordingly.
- Record the details of each newly registered patient using a form like that in Figure 2.

Using the data on the completed form you can summarise the situation and identify the individuals who delay or present with Grade 2 disability:

- Use a bar chart to summarise delay and disability grade (Figure 3).
- Look for factors such as age or sex associated with delay or disability.

Your analysis will tell you the extent of the problem and suggest groups more likely to delay.
<table>
<thead>
<tr>
<th>Card Number</th>
<th>Registration Date</th>
<th>Sex</th>
<th>Age</th>
<th>Occupation</th>
<th>Marital Status</th>
<th>Urban/Rural</th>
<th>Symptoms at Registration</th>
<th>Leprosy Group</th>
<th>WHO Grade</th>
<th>Delay in Presentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Example: ABC123</td>
<td>12/5/99</td>
<td>Male</td>
<td>35</td>
<td>Labour</td>
<td>Married</td>
<td>Rural</td>
<td>Many Patches</td>
<td>MB</td>
<td>0</td>
<td>15 months</td>
</tr>
</tbody>
</table>

Figure 2: Form for data collection from clinic records
In this data set only 2% of patients with short delays had Grade 2 disability.
In the longest delay group (more than 5 years) the rate was 14%. 

Figure 3: Example of a Bar Chart
If you understand the decisions and decision-making that result in delay then you may find a way to reduce delay. You need to find out who is involved in decision-making, the decisions that are taken, and the time committed to each action. You can then identify the main actions adding to delay and the events that finally caused the person to present and start treatment. The most direct way to do this is to listen to the experiences of individual patients and learn from them.

Q. What method will I use?

- The first method we recommend is patient interviews, using experienced staff as interviewers.
- At a later stage you might use focus groups and work with groups of patients. You might also observe the reaction of new patients to the leprosy diagnosis. (See Annex 4).

Q. How can I be sure to collect all the information I need?

- We recommend semi-structured interviews – this means following an ‘Interview Outline’ which reminds you of all the information to collect in each interview (Figure 4). Figure 5 summarises other information to record.
Q. I can think of other things I would want to ask my patients – can I change the interview outline?

- Yes, you will need to do this! What you already know or what you learn in the first few interviews will help you to develop an outline that meets the needs of your local situation.
- The fieldwork is all about learning something new, so be flexible. Follow-up new ideas or unexpected actions or comments arising in the interviews.
- When you are sure the Interview Outline is correct, print some copies with Figure 5 on the reverse. The interviewers can then use one sheet of paper for each interview and record their notes and comments from the interview on the same paper.

Q. So how do I organise the interviews?

- Make sure each patient is willing to participate. Limit interviews to 30 minutes.
- Build trust and confidence. Be ready to care for anyone who finds telling their story an emotional experience.
- Start each interview by asking the patient to describe his/her experiences.
- Ask extra questions to fill in details omitted from the patient’s story.
- Ask specific questions from the Interview Outline and other information (Figures 4 and 5).
- Finally, complete the summary and record comments about the patient and the interview (Figure 6).
Q. How should I select and train interviewers?

- Select experienced male and female staff as interviewers. A sympathetic and respectful attitude towards patients is the essential qualification.
- A male interviewer should interview male patients while a female interviewer makes notes. The roles will be reversed with female patients.
- Explain the importance of open questions and the need to learn from patients.
- Arrange for practice in interviewing, first in a role-play and later with patients.
- Start formal interviews only when you are convinced the interviewers are producing reliable data.
- Use the training interviews to further develop the Interview Outline to cover local issues.

Q. How many people do I interview?

- Plan to interview around 20 patients over a 6 week period.
- Choose patients currently receiving treatment – months 3 to 9. The experience of older patients may no longer be relevant.
- Plan a quota of interviews within age and sex groups (or other appropriate factors).
- Maintain a record of progress towards the quotas.
- You might like to match the patients for interview – one with impairment matched with one without. Later, the analysis will then focus on why the two patients had such different experience.
Looking after the data

- Leave enough time each day to complete a written interview summary for each interview.
- Don’t limit the summary to simple answers to questions. Include personal information about each patient - attitude, confidence, understanding and response to their experiences. All these are important.
- Record the patient’s experiences and comments using their own words.
- In the interview summary, clearly separate description from personal comments, interpretation and judgements.
- Make sure the interview summaries are safely filed at the end of each day.
- In a separate research diary, record the date of each interview and any comments about how the work is going or new directions that it might take.

Difficulties to expect

- Patients find it difficult to estimate time spent on individual actions. Comparison with ages of children, marriages, annual festivals etc, can help to improve the estimate.
- Patients may have difficulty in remembering where and when they looked for help. The important point is to identify the most important reason for time lost, whether through taking no action, through traditional medicine or through the health services.
- Look out for reported problems not related to leprosy. For example, there are many reasons for divorce.
- Rather than talking in general about stigma patients may describe hatred, rejection, humiliation, or people “talking bad”.
### INTERVIEW OUTLINE

**The Patient's Story:** Ask the patient to tell their story, describing each help-seeking action in detail:
- What were the symptoms?
- How important did they seem?
- Who knew of the symptoms?
- Who was involved in deciding what to do?
- What did you decide to do?
- What was the result?
- Did anyone suspect leprosy?
- How long waiting before next action?

**Describe the final decisions resulting in referral and start of treatment:**
- Who was involved?
- Who first suspected leprosy?
- How had symptoms changed by the time treatment started?
- Why did the patient finally go for treatment?

**What does the patient say was the main cause of his/her delay?**

**Before the start of treatment...**
- What did the patient believe about the cause of leprosy?
- What did they think was the cause of leprosy?

**Before the start of treatment, what did the patient think were the consequences of leprosy:**
1. Physically?
2. For general health?
3. For marriage?
4. For family members?
5. With neighbours?
6. For work or employment?

**What experience with traditional health services?**

**What experience with public/private health services?**

**What else is important** in what the patient says or what they experienced?

---

**Figure 4: Draft Interview Outline**
<table>
<thead>
<tr>
<th>Name: ____________</th>
<th>Village: ______</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinic Card Number: ______</td>
<td></td>
</tr>
<tr>
<td>Age: ____________</td>
<td>Sex: ___</td>
</tr>
<tr>
<td>Interviewed by: __________________</td>
<td></td>
</tr>
<tr>
<td>Interview Number: ____________</td>
<td></td>
</tr>
<tr>
<td>Interview Date: ______________</td>
<td></td>
</tr>
<tr>
<td>Single/Married/Divorced/Widowed: ____________</td>
<td></td>
</tr>
<tr>
<td>Education to class: ___</td>
<td>Occupation: ___</td>
</tr>
<tr>
<td>Diagnostic group (MB/PB): ___</td>
<td>WHO Grade: ___</td>
</tr>
<tr>
<td>Duration of symptoms (from notes): _____ months</td>
<td></td>
</tr>
</tbody>
</table>

**Interview Summary:**

**Interviewer’s assessment:**

What do you think was the most important cause of delay for this patient:

Write down the most important words the patient used:
(for example, hatred, isolation, fear, rejection, anger, concealment, denial, ignorance of symptoms or cure)

**Figure 5: Other information to record from interviews**
An interview summary from Malawi

Profile:
Young boy, aged 10 years, walks 16 km to clinic. PB leprosy. No previous contact. 3 months MDT completed. Anaesthetic patches. No motor or sensory loss. No ulcers.

Details:
His family members know he has leprosy. They did not reject him, neither did the village community. He can still eat with them and share their drinking water. He still contributes to the fishing nets, showing there has been little or no social impact. His first sign was an ordinary patch on his head. After a month his mother noticed and they went for traditional medicine (6 months). The traditional medicine cured the patch, but it reappeared. They did not think about going to hospital. He was identified as having leprosy by a village survey.

The parents think that everyone was affected by “magic”, women more than men, and that everyone goes to the (traditional healer) though women find it difficult because of the long distances. With leprosy they thought women were more likely to be divorced than men, because men will want to marry another woman without leprosy. They say traditional healers do not refer patients to hospital and that they can’t cure leprosy.

Comment of interviewer:
No evidence of fear or actual stigma, though there is a risk of divorce. Opting for traditional medicine caused six months delay. Traditional healers still have a strong hold over some people.

Figure 6: Interview Summary from Malawi
Chapter 6 – Learning from the community

The community’s beliefs and behaviour towards leprosy provide the background for the behaviour of patients. It is important to confirm what you learned from patients by listening to what members of the community have to say. To do this you might choose to interview community representatives or you might choose a different approach, working with selected groups of individuals.

Q. How do I choose who to speak to in the community?

First of all, choose to speak to people that have a good knowledge of how the community functions – political or religious leaders or other respected people in the community. Local teachers might be another source, though those that have moved into the area may have little local knowledge.

If you want to learn more about the importance of traditional medicine you might choose to meet with local practitioners, or with people who seek help from them.

In cities where there are medical colleges or nurse training centres you might be able to meet with new students and learn from their attitudes towards leprosy.
Q. What do I want to learn from the community?

The focus will be on key points relating to leprosy:

- What do people believe about its cause and the people it affects?
- What do people fear about the disease and the people it affects?
- Within marriages, how do people behave towards people affected by leprosy?
- Within families, how do people behave towards people affected by leprosy?
- Amongst neighbours, how do people behave towards people affected by leprosy?
- In the community, how do people behave towards people affected by leprosy?
- Are these attitudes and behaviours changing? If so, why?
- What are the most important reasons for people delaying the start of treatment?
Q. What method will I use?

- You may choose to use semi-structured interviews – but be ready to make substantial changes to the interview outline in Figure 4.
- You may choose to run Focus Groups. See Chapter 7.
- You may choose to use a method called free-listing described here (Figures 7 and 8).
- Following your free-listing with community representatives you might plan a free-listing exercise with patients or with their family members.

Q. How does free-listing work?

In free-listing you ask people to write down how they respond to one or more carefully chosen questions. For example, you might ask people to write down what they think in response to one of the questions following:

- For me to be diagnosed with leprosy would mean …
- Telling my husband (wife) I had leprosy would mean …
- Telling my neighbours I had leprosy would mean …
- The reasons I delayed starting my treatment for leprosy were …

Personalising the questions encourages people to express their attitudes and fears and reveals their understanding of the underlying issues.
Organising a free-listing exercise

- Get participants seated and provide them with pen and paper.
- Where literacy is a problem, form groups and ask one person to act as secretary.
- Write the chosen question on a board for all to see.
- Explain that there will be 10 minutes to think and write down responses to the question.
- Make it clear that there are no right or wrong answers! People should simply write what they think. Each person may write several different responses.

- After 10 minutes, go into a group session, asking participants to report their responses.
- Write the responses on the board, counting how many people give each response.
- Identify any groupings – e.g. positive or negative attitudes, knowledge or ignorance of the disease, fears for physical consequences, fears for social consequences, impact on family, impact on work.
- Identify the most frequent and most important responses.
- Explore these through group discussion.
- Record all the responses and make notes from the discussion for inclusion in the final analysis and report.
Free-listing with nurse trainees in India

The exercise took place in a leprosy hospital situated in a city centre in India. It was the first action at the start of a two-day orientation course for nurse trainees from a local training centre.

- The trainees are asked to write down how they would respond if told they had leprosy.
- The trainees wrote down their responses.
- The responses were written onto a blackboard, counted and discussed.

The response in 2001 ranged from complete confidence in the cure and in the response of family members to complete despair – I want to die. The trainees recognised the potential social implications – people may not touch me or love me – but there is also the potential for self-imposed isolation – shall I practice homosegregation.

The results make it clear that despite their educational attainments the nurses come from a culture where fear of leprosy persists. There is a substantial risk of some form of adverse social impact. Leprosy also has a significant psychological impact – the fear of what may happen. Patients at the same centre confirmed that social pressures relating to leprosy affected their decision-making.

Figure 7: Free-listing with nurse trainees in India
For me to be diagnosed with leprosy would mean …
Responses of 39 nurse-trainees at the start of a two-day orientation course in leprosy.

1. I am uncertain how society will treat me (20)
2. I would doubt if my near ones would accept me (16)
3. How will I live in the world? (13)
4. Why did I have to get this disease? (11)
5. I want to leave my home (9)
6. Who will marry me (8)
7. Should I practice segregation at home? (8)
8. I want to die (7)
9. Will I be treated the same was as others in society? (7)
10. How will I attend my college? (4)
11. Will my children get leprosy? (4)

All the trainees were aware of the cure. Smaller numbers of responses included a curse from God, it relates to past sins, people will not touch me, I will have to depend on others, I feel like an orphan.

Figure 8: Result of a free-listing exercise in an Indian city in 2001.
Chapter 7. Learning from staff

The experience of your staff is as important as the knowledge of the community in validating the experiences of your patients. Staff members meet with patients every working day. They listen to patients and try to understand their symptoms and experiences. Creating the opportunity to learn from staff adds value to the staff and provides insights into their understanding of what contributes to delay.

- Many staff have an intimate knowledge of the local culture.
- The best staff have good working relationships with patients.
- They know where patients live and may be aware of the home situation.

To collect information from staff you might choose to interview selected staff members, you might choose a free-listing exercise or you might choose to use focus groups, as described here.

Through your group work with staff you might identify further information to collect from patients. You might add to the Interview Outline or plan group work with patients.

Figures 9 and 10 describe group work with staff in Paraguay.
Q. How do I organise a focus group?

Arrange for the group to meet together and encourage them to discuss the reasons why patients delay.

- Decide on the focus - no more than one or two questions for discussion – e.g. what contributed to your delay?
- Plan the meeting in advance – the time and place. Choose carefully who to invite, covering appropriate groupings by sex, age or other important variables.
- Write guidelines to ensure that the discussion proceeds in an orderly fashion. Appoint and instruct a facilitator who should:
  - Strictly limit their involvement in the discussion.
  - Ask only an occasional question that ensures the discussion remains focused.
  - Try to be just a “fly on the wall”, listening and learning from what is said.
- An observer should take notes, leaving the facilitator free to concentrate on the direction of the discussion.
- You might decide to record the discussion, but this can be difficult amongst many people in a large room.
- Set a strict limit on the time of the meeting – e.g. 45 to 60 minutes.
- The observer and facilitator need to agree and complete a written summary of each group meeting, including conclusions of those present. These are the basis for analysis using the method described in Chapter 8.
- Note the points of agreement and disagreement.
- Look out for experiences and information that are of most interest.
- Finally, ask the facilitator to identify the primary factors contributing to delay.
Practical points for group work:

- Expect the unexpected – unforeseen ideas will arise.
- Try to identify any important differences between group members.
- Interactions between group members will develop. There may be points of special interest or points of strong emotional involvement.
- Contradictory opinions and feelings may be expressed.
- How well the group work is facilitated strongly influences the result.
- Encourage everyone in the group to participate.

Q. Need more information?

- Repeat the exercise with patients, discussing issues relating to the impact of the diagnosis or the things people fear about leprosy.
- Try group work with representatives of the community (e.g. teachers or religious leaders) or with patients and their family members.
Group work with staff in Paraguay

The exercise took place in the main referral hospital in Paraguay. Many of the staff have been involved in the work for 20 years or more. In addition to in-patient and out-patient work at the centre there are out-patient clinics linked to government health centres across the country.

Discussions focussed on:

- Local knowledge and awareness of leprosy
- The reasons for delay
- Any changes taking place

There was enthusiastic discussion that opened up subjects related to attitude and behaviour towards leprosy and what might be addressed through health education. We also interviewed two senior staff of the project.

We found a high level of agreement between the information provided by staff and the information from patient interviews.

Figure 9: Group work with staff in Paraguay
Results of group work with staff in Paraguay

Discussion on “Why people delay in starting treatment”. The staff were organised into two groups of 8.

The groups agreed that the most important reasons for delay were ignorance and failure to recognise the importance of symptoms. Next, female staff identified use of traditional medicine, the male staff identified wrong diagnoses made by doctors in the private and general health services. Male staff also identified a cause of delay relating to culture, which defines or requires serious disease to be accompanied by pain. Since leprosy causes no pain it is not recognised as a serious disease, so people do not seek treatment. The groups also identified other contributors to delay: fear of the diagnosis, the perceived cost of the treatment, fatalism and denial.

Special needs identified included: people living in isolated areas, places where witchdoctors have power and forbid contact with the health services, and people experiencing no pain.

The groups identified necessary actions - improved communication about leprosy, increased awareness of the cure, greater awareness and better diagnostic skills in the health service.

Figure 10: Summary of focus group discussion with school teachers in Paraguay
Chapter 8  Analysing your data

Your fieldwork has produced valuable data. You now need to identify the most important findings and write a short report. This is potentially a complicated area, with the risk that you get submerged in detail. The key to success will be to use the simplest possible approach, as described here.

Because the data is in text form the analysis involves reading and rereading and preparing a written summary.

- First check you have all the data, from patient records, from interview and from group work.
- Read through all the data 2 or 3 times:
  - Identify the main two or three actions that contribute to early or late presentation.
  - Identify the one or two groups of individuals that are likely to delay.
  - Identify the one or two geographical locations from which people are likely to delay.
- Prepare a summary table (Figure 11) to compare what was said by patients, by community representatives and by staff.
- If you don’t believe what you read or find information is missing, go back to patients, staff or community to fill the gaps.
- Read through the data one final time to make sure nothing has been missed

Finally:

- Identify the most important causes of delay confirmed by at least two sources.
- Ask yourself what these findings mean for the project.
Q. What do I write in my report?

In writing your report answer the questions:

- What are the most common reasons for delay?
- Who are the people most at risk?
- What is the extent of the problem?
- Make recommendations about what might be done about it.

Remember:

- Keep the report short, no more than 4 pages.
- Focus on the main findings.
- Include a Summary and Conclusions
- Identify recommended actions to be taken in response to your findings (Chapter 9).
- Write and circulate the report within two weeks of finishing fieldwork.
<table>
<thead>
<tr>
<th>Sources:</th>
<th>Clinic records</th>
<th>Patients</th>
<th>Community</th>
<th>Staff</th>
</tr>
</thead>
<tbody>
<tr>
<td>Causes of delay:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>People at risk of delay:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Locations where delay is most common:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 11: Summary chart for analysis
Chapter 9 – Applying your results

- When you review your findings you might ask yourself three questions:
  - Is there something new we should be doing?
  - Is there something already started that we should be doing better?
  - Is there something already started that we should do less?

- Remember that your fieldwork is limited, so be cautious in recommending change. After 6 months repeat some of the fieldwork to see what the effect of the changes has been.

- With your senior manager you might review your findings alongside a summary of your project’s objectives and activities. The following SWOT analysis could be used (Figure 12).
**Figure 12: Summary of conclusions and resulting application**

<table>
<thead>
<tr>
<th>Findings:</th>
<th>Application:</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strengths:</strong></td>
<td><strong>Management:</strong></td>
</tr>
<tr>
<td>Good awareness in urban area.</td>
<td>Level of respect affirms effectiveness of project over long period.</td>
</tr>
<tr>
<td>Effective lay referral.</td>
<td>Consider stopping the labelling of symptoms or disease as leprosy.</td>
</tr>
<tr>
<td>Good referral from health services.</td>
<td></td>
</tr>
<tr>
<td><strong>Weaknesses:</strong></td>
<td></td>
</tr>
<tr>
<td>Preference for alternative medicine.</td>
<td></td>
</tr>
<tr>
<td>Variable referral practice by private doctors.</td>
<td></td>
</tr>
<tr>
<td>Need more information on why individuals delay.</td>
<td></td>
</tr>
<tr>
<td><strong>Opportunities:</strong></td>
<td><strong>Programme:</strong></td>
</tr>
<tr>
<td>Able to train and encourage alternative practitioners to recognise leprosy and refer?</td>
<td>Need for effective health education in the community at large and with alternative practitioners to address the persistence of traditional beliefs.</td>
</tr>
<tr>
<td>Opportunities to access electronic and printed media.</td>
<td>Need to improve communication with health services staff.</td>
</tr>
<tr>
<td><strong>Threats:</strong></td>
<td>Need to develop contacts with and knowledge of private doctors.</td>
</tr>
<tr>
<td>Persistence of traditional fears of physical and social consequences of leprosy.</td>
<td>Identify ways to address psychological impact and fears associated with diagnosis.</td>
</tr>
<tr>
<td></td>
<td>Teach staff some basic counselling skills?</td>
</tr>
</tbody>
</table>
Application

Need more information?

If you still need more information on the causes of delay you might add questions to the forms completed with each newly registered patient (Figure 13).

| Duration of symptoms (months): |
| Delay before first action (months): |
| How much time did the patient spend in seeking help from other sources: |
| Traditional practitioners: _____ (months) | Village doctors: _____ (months) |
| Homeopathic practitioners: _____ (months) | Medicine Shops: _____ (months) |
| Ayurvedic practitioners: _____ (months) | Private Doctors: _____ (months) |
| Natural practitioners: _____ (months) | Health Service Doctors: _____ (months) |
| Other practitioners: _____ (months) |  |

Figure 13: Information to collect from newly registered patients
Annex 1 - Persistent stigma

In 1989 Dennis Mull wrote this about the history of stigma related to leprosy:

*Throughout recorded history, whether in Asia, Europe, Africa or the New World, leprosy victims have been shunned and in a very real sense discarded by the societies in which they live. Such behaviours arise from a belief that the disease is a punishment from God, with strong associations that only the most heinous sins would have earned so fearful a punishment. People were caste out of their homes as a threat to society.*

(Mull JD et al, Social Science and Medicine, 29(7) 1989, 799-811.)

The research that resulted in the production of this booklet has demonstrated that the same attitudes and behaviour exist at the beginning of the 21st Century.
Annex 2 – Fieldwork Skills

Your interviewers need to be able to encourage people to speak freely and to tell their story. They need analytical skills to understand how patients decided to seek help and the sequence of actions they took. They must be able to identify the most important features of each individual’s experience.

The basic skill requirements include:

- Experience in basic interview technique.
- Experience in organising and leading group discussions.
- A sympathetic and caring attitude towards patients.
- Analytical skill, able to organise and record the experiences of each individual from often muddled narrative.

You should consider:

- The special needs of women.
- The special needs of children - probably addressed through an accompanying adult or older sibling.
- The potential role for a senior worker as a fieldwork supervisor.
What makes a good interviewer/facilitator?

- Keep to the interview/group work outline and be sure to cover all the questions.
- Use open questions, those that encourage the patient to talk freely and confidently:

<table>
<thead>
<tr>
<th>Open questions encourage detailed answers</th>
<th>Closed questions invite one word answers</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Example:</strong></td>
<td><strong>Example:</strong></td>
</tr>
<tr>
<td>Tell me about your first symptom</td>
<td>Was your first symptom a patch?</td>
</tr>
</tbody>
</table>

- Work to develop good listening skills:
  - Repeat words and phrases used by the patient – this encourages them to explain or give more detail.
  - Provide a private area for the interview, free from interruptions.
  - Sit alongside rather than facing across a desk.
  - Respect the point of view of the patient. Do not be judgmental.
  - Maintain an open mind – be ready to learn something new.
  - Be sensitive. For many, leprosy is a major crisis with emotional, psychological or social impact.
Annex 3 - Making an Action Plan

Preparing an Action Plan will require you to consider all the details of how and when you want to undertake fieldwork. Include times to meet with supervisors and time to deal with problems that might arise.

You may need to consider the following:

- Do you need organisational approval?
- Do you need ethical approval?
- Are there any special requirements relating to your integrated programmes?

Plan the work in phases - preparation, data collection and reporting, each with definite start and end dates.

Phase I – Preparation – weeks 1 - 2

- Take two weeks to set objectives and complete planning.
- Select staff to train for interview and other group work.
- Prepare paperwork to record data.
- Plan the selection of patients, staff, community representatives etc to interview.
- Schedule interviews, group work and other fieldwork.
- Keep costs to a minimum by working through scheduled clinics.
- Complete training.
- Finalise Interview Outlines, Focus Group and Free-listing questions.
Phase II – Data collection – weeks 3 – 8

- Extract data from patient records.
- Work at completing a weekly quota of patient interviews plus community and staff interviews or group work.
- During the period, plan at least three meetings to review progress and solve problems.
- Monitor the effectiveness of the fieldwork and be ready to change the focus if necessary.
- Check records of each interview and group meeting are adequate.
- Check data is stored safely.

Phase III - Analysis and reporting – weeks 8 - 9

- Collate data, checking that all fieldwork is complete.
- Summarise the data.
- Prepare a written report, including recommendations.
- Prepare a statement of costs.
- Communicate your findings
- Begin work at their application.
Annex 4 - Choosing Methods

- **Semi-structured interviews** - allow you to encourage individual patients to share their experiences while you follow an Interview Outline to ensure you collect all the information you need. The interviewer may be flexible in using the Interview Outline, but must always be careful to cover all the listed topics.

- **Focus Groups** – provide the opportunity for a small number of people to discuss a topic freely and spontaneously under the guidance of a facilitator. They are especially helpful for gaining insight into people’s perceptions, attitudes, opinions, behaviour and experiences. They can also help to assess the acceptance of concepts and how information is being passed on from one person to the next.

- **Free-listing** – a simple method to gather knowledge about specific categories or items of information.
• **Observation.** Recommended as way to assess the effectiveness of the interview process. It might be used to assess the impact of the leprosy diagnosis. As the name suggests, it involves watching behaviour and assessing the changes that occur over time. In Paraguay we observed the traumatic impact of the leprosy diagnosis on two young women. Elsewhere we relied on observation to assess the emotional and organisational state of patients.

• **Survey.** We recommend you don’t rely on this approach. More than likely it will only confirm what you already know. Surveys need comparatively large numbers and a complicated analysis.

**Other points about fieldwork**

Within the constraints of time and resources we recommend you make use of all available sources of information. In addition to patients, community representatives and staff you may choose to work with health service practitioners, traditional healers, professionals in health education, former patients and the family members of patients.

Such alternative sources, plus historical records and reports will add to the data available to you in analysis and report writing.
Annex 5 - Evaluation

Evaluation - I

We recommend you evaluate the results of your fieldwork. First you might assess their value by reviewing your findings. How does the value of what you learned compare with the effort put into the fieldwork?

After twelve months you might complete a more detailed evaluation. What has been the impact on the project?

Evaluation - II

As part of our evaluation of the procedures described here, we would be interested to hear you assessment. Please forward comments to the contact address given at the front of the booklet.

We have four main areas of interest:

- Please describe any difficulty you experienced in understanding or implementing fieldwork.
- What did you learn from the fieldwork?
- What additional methods or sources did you use?
- What has been the impact on your work?