Module: Epilepsy

Overview

Learning objectives

• Promote respect and dignity for people with epilepsy.
• Know common presentations of epilepsy.
• Know the assessment principles of epilepsy.
• Use effective communication skills in interactions with people with epilepsy.
• Know the management principles of epilepsy.
• Perform an assessment for epilepsy.
• Assess and manage physical health in epilepsy.
• Assess and manage emergency presentations of epilepsy.
• Provide psychosocial interventions to persons with epilepsy and their carers.
• Deliver pharmacological interventions as needed and appropriate in epilepsy considering special populations.
• Plan and perform follow-up for epilepsy.
• Refer to specialists and link with outside agencies for epilepsy as appropriate and available.

Key messages

• Epilepsy is not inherited or contagious.
• Assessment includes:
  – Assessing and managing an acute/emergency presentation.
  – Assessing for epilepsy and any other underlying causes of the seizures.
• Seizures are symptoms and not the cause, therefore underlying causes should always be explored and assessed.
• To be considered epileptic there must be two or more unprovoked, recurrent seizures.
• Epilepsy can be treated effectively with antiepileptic drugs in non-specialized health settings.
• Psychoeducation and psychosocial interventions to promote functioning in daily activities are empowering for the person with epilepsy to enable them to manage their condition.
• Adherence to treatment and regular follow-up are critical.
• People with epilepsy can lead normal lives.
• Children with epilepsy can go to a normal school.
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**Total duration (without breaks) = 4 hours 30 minutes**
Session 1. Introduction to epilepsy

40 minutes

Session outline

- Introduction to epilepsy.
- Assessment of epilepsy.
- Management of epilepsy.
- Follow-up of a person with epilepsy.
- Review or materials and skills.

Begin the session by briefly listing the topics that will be covered.

Activity 1: Person’s story followed by group discussion

Using the person’s story to:
- Introduce the activity and ensure participants have access to pens and paper.
- Tell the person's story – be creative in how you tell the story to ensure the participants are engaged.
- Immediate first thoughts – give participants time to give their immediate reflections on the story.

Ask participants to think about people they have cared for in the past with epilepsy? Can they think of any cases? How did the person with epilepsy behave, how did their family and carers cope?
Write a list of local terms and descriptions for epilepsy and compare those with common presentations described in the mhGAP-IG.

(Maximum five minutes.)

Read through the common presentations of people with epilepsy.

Talk through the points on the slide by explaining that epilepsy is a neurological condition characterized by recurrent seizures.

Seizures are brief disturbances in the electrical functions of the brain.

There are potentially many different causes of epilepsy but it is not always easy to identify one.

Talk through the possible causes.
Explain the signs and symptoms of epilepsy. It is typified by seizures.

In order to receive a diagnosis of epilepsy, there needs to have been two or more recurrent unprovoked seizures (in the past 12 months):
- Recurrent = usually separated by days, weeks or months.
- Unprovoked = there is no evidence of an acute cause of the seizure (e.g. febrile seizure in a young child).

Seizures are brief disturbances of the electrical function of the brain. Characteristics of seizures vary and depend on where in the brain the disturbances first start and how far it spreads.

Describe the two types of epilepsy as described on the slide.

Explain that this module will focus on convulsive epilepsy, as that is the type associated with more fear, stigma and discrimination.

Talk through the points on the slide and briefly explain what a seizure is.

Highlight again that in this module we will concentrate on convulsive seizures as 70% of all seizures are convulsive.

Convulsive seizures have a high mortality rate, but they can be treated.
Use the slide to explain:
• What a person is likely to experience during a seizure.
• What the person is likely to experience after the seizure.

Explain that epilepsy is not contagious.

Talk through the points on the slide.

Facilitate a brief discussion about which of these conditions is a common cause of epilepsy in their local community.

It is important to know and discuss local environmental factors that could contribute to seizures and epilepsy.

Encourage participants to participate in the discussion to make sure they are aware of the local causes.

Emphasize the first point on the slide indicating that epilepsy can be treated effectively in non-specialized health settings.

When people are treated they have a good prognosis. Two to five years’ successful treatment and being seizure-free means medication can be stopped in 70% of children and 60% of adults.

Antiepileptic medication is affordable – US$ 5 per year.

In low- and middle-income countries about 75% of people with epilepsy may not receive the treatment they need.

In fact, in low- and middle-income countries there is a low availability of antiepileptic drugs (AEDs) – this may act as a barrier to accessing treatment.
Here are some other reasons (although not exhaustive) for the high treatment gap:

- Epilepsy is a low priority for many countries.
- Limited capacity of health-care systems to address epilepsy and inequitable distribution of resources.
- Lack or severe shortage of appropriately trained staff.
- Inadequate and inconsistent access to affordable medicines.
- Societal misconceptions.
- Poverty.

Generate a brief discussion. Revisit the list of local names and terms produced for a person with epilepsy.

Ask the group if some of the names and terms are negative?

How might that make the person/family feel?

How might that impact on their likelihood to seek help?

Explain that people living with epilepsy around the world are quite often stigmatized and discriminated against.

Common misconceptions about epilepsy are that it is contagious, and people must be avoided and feared; and that they are possessed by evil spirits and/or bad in some way.

People are denied access to health care and treatment, or they are too afraid to seek help.

Often children are withdrawn from schools. People with epilepsy are overlooked for jobs (impacting on their ability to earn money and support themselves and their family). People with epilepsy are often unable to get married and sometimes prevented from driving.

To summarize, even though epilepsy is a very treatable condition, people with epilepsy are not receiving the help they need and instead are being stigmatized and discriminated against.

**Local names for epilepsy**

- Are the names/local descriptions of epilepsy negative?
  - Some of the local terms may imply a person is mad, possessed, stupid or cursed.
  - How might this impact on a person and their family?
  - How might this impact on their likelihood to seek help?
Approximately 50 million people worldwide have epilepsy, making it one of the most common neurological diseases globally.

Nearly 80% of the people with epilepsy live in low- and middle-income countries.

People with epilepsy respond to treatment approximately 70% of the time.

Nearly 75% of people with epilepsy living in low- and middle-income countries do not get the treatment they need. In some regions of the world, like Africa, this can be as high as 85%.

Those with epilepsy have a three to six times greater risk of dying prematurely but epilepsy can be treated effectively in primary health care.

Treatment is simple, inexpensive and effective. Some 70% can be seizure-free for life after two years of treatment.
Session 2.
Assessment of epilepsy

1 hour 40 minutes

The first part of the session focuses on the management of acute seizures and emergency presentations. The second half of the session focuses on how to assess someone for epilepsy.

Explain that there are two ways that people with epilepsy enter health care services:
• During a seizure – as an emergency presentation.
• After a seizure.

Have participants read through the assessment principles for epilepsy.

mhGAP-IG has an assessment algorithm for both and in this training we will start with how to manage seizures which present as emergencies.

Emphasize why managing seizures is an emergency.

Talk through the points on the slide.

**Why are seizures treated as an emergency?**

- Treatment can end seizures or shorten seizure duration, which limits the damage they can cause.
- Prolonged or repeated seizures can result in brain injury.
- Prolonged or repeated seizures can result in death if not treated immediately.
- Seizures can be a symptom of a life threatening problem, like meningitis.
Activity 2: Group discussion: Emergency presentations

A person is brought into the clinic and is unconscious after a reported seizure.

What are your first actions?

Duration: 10 minutes.

Purpose: To learn how much participants know about managing acute seizures.

Instructions:
- Give individuals a few minutes to think individually about what they would do in this situation.
- Facilitate a group discussion and seek group consensus to create a comprehensive list of steps they would take to help the person.

Talk through steps 1 and 2 in the algorithm as shown on page 53 of mhGAP-IG Version 2.0.

Emphasize that participants need to understand that they cannot wait until they establish a complete diagnosis to start managing the seizure. Management and assessment must happen at the same time.

Ensuring the A, B, C (airways, breathing, circulation) is crucial, even if they do not have a clear idea about the cause yet.
Check with participants if they’ve already had training on this topic.

If this is a new topic then ensure you give sufficient time to ensure participants understand how to manage acute seizures.

If they have received training in this then explain that this is an opportunity for them to refresh their knowledge.

Ask participants to explain and then demonstrate how they put a person in the recovery position (20 minutes).

Divide the participants into pairs and have them practise putting each other into the recovery position (15 minutes).

**Recovery position**

A. Kneel on the floor to one side of the person. Place the person's arm that is nearest you at a right angle to their body, so it is bent at the elbow with the hand pointing upwards. This will keep it out of the way when you roll them over.

B. Gently pick up their other hand with your palm against theirs (palm to palm). Now place the back of their hand onto their opposite cheek (for example, against their left cheek if it is their right hand).

C. Now use your other arm to reach across to the person's knee that is furthest from you, and pull it up so that their leg is bent and their foot is flat on the floor.

D. Now, with your hand still on the person's knee, pull their knee towards you so they roll over onto their side, facing you.
Emphasize that these vital signs need to be measured and documented.

Respiratory rate actually needs to be counted, not estimated, since trends in respiratory rate become quite important if the person has recurrent seizures and requires aggressive treatment with multiple doses of medications, which can suppress the respiratory drive.

- Time the duration of the convulsions.
- Make sure the person is in a safe place – ensure that nothing is likely to fall on them and/or they can’t hit anything if they convulse.
- If possible place in an i.v. line for medication/fluids.
- Know when to refer – if a person has a head injury, a neuroinfection or focal neurological deficits then refer to hospital.

Direct participants to page 54 of mhGAP-IG Version 2.0.

Talk through the next steps highlighting the special population: pregnancy/post-partum and when to suspect eclampsia.

A pregnant woman who has no history of epilepsy and presents with seizures may have eclampsia.

Eclampsia is a condition in which one or more convulsions occur in a pregnant woman suffering from high blood pressure.

The condition poses a threat to the health of the mother and the baby.

If there is a midwife in your clinic call them to assist. They may have training in how to support people with eclampsia.

Refer immediately to a hospital.
Step 4 (give medication to stop convulsion) – if you cannot establish an i.v., **do not give diazepam intramuscularly** (i.m.). Ask participants if they know why they should not give i.m. diazepam?

Explain that i.m. diazepam is poorly and unpredictably absorbed and diazepam should only be given rectally.

Explain how to give rectal diazepam. Mention to participants that they need to teach this to the carers of people with seizures for them to be able to do it at home.

**Instructions:**
- Draw up the dose from an ampoule of diazepam into a tuberculin (1 ml) syringe.
- Base the dose on the weight of the child, where possible.
- Remove the needle.
- Insert the syringe into the rectum 4 to 5 cms and inject the diazepam solution.
- Hold buttocks together for a few minutes.
- If the convulsion continues after 10 minutes, give a second dose of diazepam rectally (or give diazepam intravenously (0.05 ml/kg) if i.v. infusion is running.

Ask participants what should they do if the convulsions have not stopped within 10 minutes of the first dose of medication?

Direct the participants to page 55 mhGAP-IG Version 2.0. Talk through steps 5 and 6.

Explain that we will look at the management protocols in the next session but for now we will concentrate on the assessments.

Ask participants if they know when a person is in status epilepticus?
Direct the participants to page 56 mhGAP-IG Version 2.0.

- Explain the definition of status epilepticus and emphasize that management should occur in a health facility.
- Continue talking through the steps of the algorithm if a person is in status epilepticus.
- Explain that i.v. antiepileptic medicines such as i.v. phenytoin and phenobarbital should always be administered in a health care setting.

Once the convulsions have stopped, take step 10 (page 57 mhGAP-IG) – evaluate (and treat as appropriate) for underlying cause of convulsions.

Remind participants that seizures are symptoms not causes, so you always need to look for the cause.

If the person presents convulsing, it is an emergency and needs to be treated urgently as:

- Seizures can be a sign of a life-threatening problem.
- Seizures can result in brain injury or death.

Explain that we are now going to look at some possible causes in more detail.

**Underlying causes**

Explain that if you suspect a brain infection is causing the seizures, establish if there is a fever, vomiting or a rash.

If there are then manage the seizure as discussed.

Initiate treatment for underlying brain infection (such as i.v. antibiotic for meningitis).

Briefly mention specific treatments or national guidelines for common infections such as cerebral malaria, meningitis, neurocysticercosis (WHO is currently developing guidelines for the treatment of neurocysticercosis), etc.
Explain that another cause of the seizure could be trauma.

If they suspect trauma they should talk through the points on the slide.

This may be a good opportunity to ask participants which methods they might use to stabilize the neck.

If there seems to be some confusion or inappropriate ideas, find a volunteer and show them with a hands-on example.

Ask the group what they have available to stabilize the neck or what they might be able to make with local materials. The participants may be able to offer each other advice.

Ask participants how they would check for evidence of trauma before revealing the answers.

Ask participants how they would check for evidence of trauma:
• stroke and tumour
• drug ingestion or alcohol withdrawal
• metabolic abnormality.

Remind participants that we have already discussed what to do if we suspect eclampsia in pregnant or post-partum women.

But ask the participants what they would do if the person is a child with fever?

Then reveal the answers.

Febrile seizures are common in primary health-care settings.

Ask the participants to explain the difference between febrile seizures and epilepsy.
Answer: In epilepsy the person has recurrent, unprovoked seizures without fever.

Febrile seizures occur when the child has a high fever.

Make it clear that febrile seizures are not epilepsy.

Clarify that it is important to rule out complex febrile seizures as these are at a higher risk of serious underlying pathology and generally need hospital admission, CT scan and lumbar puncture.

What is a complex febrile seizure?

It is a complex febrile seizure if one of the following criteria is present:
- **Focal**: Starts in one part of the body.
- **Prolonged**: More than 15 minutes.
- **Repetitive**: More than one episode during the current illness.

A complex febrile seizure needs to be referred to hospital.

Explain the criteria for complex febrile seizures:
- **Focal**: For example, the seizure starts in the arm and then generalizes to the entire body.
- **Prolonged**: Even if it is due to fever with no signs and symptoms suggestive of neurological illness or brain infection, if the duration is more than 15 minutes, it is considered a complex seizure.
- **Repetitive**: If seizures are repetitive, it is considered to be complex.

Emphasize that you must refer the person to hospital.

Management of simple febrile seizures

1. Look for possible causes and manage fever according to the local IMCI guidelines.
2. Observe for 24 hours.
3. Follow-up in one to two months to assure no further seizures.

Talk through the points on the slide.

Simple febrile seizures usually last for less than five minutes. The child will:
- Become stiff and their arms and legs may begin to twitch.
- Lose consciousness and may wet or soil themselves.
- If there is only one seizure, it can leave the child feeling sleepy for up to an hour afterwards.

A simple febrile seizure like this will only happen once during your child’s illness.
Refer to the WHO Integrated Management of Childhood Illness (IMCI) guidelines, if needed, for more details. If the community does not use or is not aware of the IMCI guidelines, refer to: http://whqlibdoc.who.int/publications/2008/9789241597289_eng.pdf

Explain to participants that they have looked at the emergency assessment and management of acute convulsions.

Once the convulsions have stopped and the person has had sufficient time to rest and recover, the next step is to assess for epilepsy.

Activity 3: Video demonstration: Assessment

Explain to the participants that they are about to watch a video of a person being assessed for epilepsy by a primary healthcare professional.

During the video participants should look at the epilepsy assessment on page 58 of the mhGAP-IG Version 2.0 and use the clinical decision-making points to decide if the person has epilepsy.

https://www.youtube.com/watch?v=RUIRg55sx10&index=6&list=PLU4ieskOli8GicaEnDwe5Q6-yaGxhes5v.
After watching the video ask participants:
• Does Faten’s presentation match that described in the common presentation of epilepsy?
• Does Faten have convulsive seizures?

Seek a group consensus.

Ask the participants:
• Has Faten had at least two of the symptoms described during an episode?

Seek group consensus.

If yes, then suspect convulsive seizures.

Explain that the next step in the assessment is to find out if there is an acute cause?

How did the health-care provider assess if there was a neuroinfection or other possible cause?

Note: Replay the video as many times as necessary to ensure participants provide an answer.

Ask the participants:
• Does Faten have epilepsy?
• How did the health-care provider assess if Faten had had at least two seizures on two different days in the past year?
• How did the health-care provider do a physical examination? What did he look for?
What to look for on physical examination?

- Signs of head and/or spinal trauma.
- Signs of meningitis: stiff neck, vomiting.
- Weakness on one side of body or in one limb.
  - In unconscious people who are unresponsive to pain, you may notice that one limb or side of the body is “floppy” compared with the other.

Ask about other medical conditions

1. Are they diabetic? Are they on any medications?
   - Could this be low blood sugar?
2. Are they HIV positive? Are they on any medications?
   - Could this be an infection (e.g. meningitis)?
3. Is there any chance of poisoning?
4. Is this person a drug user or a heavy drinker?
   - If yes, in addition to managing their acute seizures, you will need to do an assessment according to the drug and alcohol use sections of the mhGAP-IG.

Talk through the points on the slide.

If there are conditions in the region especially likely to cause seizures, discuss these here, e.g. cerebral malaria or Japanese encephalitis, neurocysticercosis.

Talk through the points on the slide and ask these questions to try and find out about any other medical conditions that could be causing the seizure.

Mention common causes of poisonings and drugs used in the environment, e.g. in farming communities organophosphate poisoning is common.

It may be worthwhile discussing other signs and symptoms of common exposures in the environment.

This added information makes the training more relevant and may help the participants remember what you taught them next time they see such a case.

Ask the participants:
- Did the health-care provider assess for concurrent priority MNS conditions?
- Do you suspect Faten has any symptoms of different MNS conditions?
- Does Faten show any imminent risk of suicide?
Activity 4: Role play: Assessment

Activity 4: Role play

- A person comes to a primary health-care clinic for the first time after they had a fainting spell the week before.
- The person comes with their spouse.
- The health-care provider conducts an assessment using the algorithm on page 58 of the mhGAP-IG Version 2.0.

See EPI supporting material role play 1.

Print off four different instruction sheets for the participants playing different roles.

Ensure the person playing the role of the observer also has a competency assessment form (see ToHP training forms) in order to assess the participants.

Duration: 30 minutes.

Purpose: This role play enables participants to practise conducting an assessment to establish if someone has epilepsy.

Situation:
- A person comes to a non-specialized health setting for the first time after they had a fainting spell the week before.
- The person comes with their spouse.
- The health-care provider conducts an assessment using the algorithm on page 58 of the mhGAP-IG Version 2.0.

Instructions:
- Divide the participants into groups of four.
- Instruct one person to play the role of the health-care provider, one the person seeking help, one person the spouse and one the observer.
- Distribute the role play instructions to each person depending on their role.
- Ensure that the participants keep to the allotted time.
Session 3.
Management of epilepsy

1 hour 15 minutes

Begin by asking participants what management intervention strategies they think might be appropriate for people suffering with epilepsy.

Explain that if the person and the family are also experiencing high levels of discrimination and/or stress then relaxation strategies and strengthening social support strategies can also be used.
Explain that managing epilepsy with pharmacological interventions and in special populations will be discussed soon, but first psychoeducation will be considered.

Ask participants to read through page 64 mhGAP-IG Version 2.0 (psychosocial interventions).

Clarify any concerns/questions participants may have.

Group activity: In plenary, ask participants to adapt any psychoeducation messages to be culturally appropriate in the local context.

Emphasize that a seizure diary can be very helpful in managing epilepsy.

It is useful because it gives a clear idea about the person’s problems and how they are progressing.

It also empowers the individual to gain some control over their epilepsy and learn:
- When their seizures happen and what triggers them.
- How medication is having an effect on them.

Make it clear that the diary does not have to be exactly as displayed.

Any record will suffice as long as it includes the details of the event:
- Whether the person was taking the medicines regularly.
- What happened.
- When it happened.
- What/if any triggers were present.
Explain that people with epilepsy can also learn to manage their seizures and understand them better by seeking witness accounts of their seizures. Also, discussing the lead up to their seizures with carers, family members etc. can help.

**Group discussion:** First, ask participants what medications they use to manage epilepsy and discuss in the group.

Give the participants five minutes to read through point 2.3 (Initiate antiepileptic medication) and look at Table 1.

Ask participants to share what key messages they found most important?

Point out the risks of prescribing medication to special populations.

Highlight that once the appropriate medication has been chosen, ensure that it is consistently available.

- Only start one medication.
- Start at the lowest dose.
- “Go slow”, increase the dose slowly until convulsions are controlled.
- Consider monitoring blood count, blood chemistry and liver function, if available.

Remind participants of the instruction in the mhGAP-IG to choose a medication that will be **consistently available**. Ask them to reflect whether that is realistic in their settings?

**Facilitate a discussion about:**
- What drugs are available in your setting?
- How much does the medication cost?
- How can you ensure medication adherence?
- What can you do if the medication is not consistently available?
Psychoeducation for medication management

Explain to the person and the family:
• The need for prompt medical treatment.
• Explain that this is a chronic condition and the medication must be taken as prescribed.
• If you take the medication as prescribed then the majority of people find that the seizures are fully controlled.
• Explain the potential side-effects and what to do if they occur.
• Explain the risk of further seizures if doses are missed.
• Plan for regular follow-ups.

Psychoeducation for medication management
Talk through the points on the slide and use the below for extra emphasis.

Key messages:
• Explain to the person and the carer the need for medication.
• Explain the importance of taking the medication as prescribed.
• Explain that if they take the medication as prescribed they can expect to control the seizures.
• Explain the potential side-effects and what to look out for and what to do.
• Explain the risk of further seizures if doses are missed.
• Plan for a follow-up session to show that you are still there to support them.

Ask participants to read through the management options for special populations.

Ask participants:
• Why these groups are considered special populations?
• What are the concerns for:
  – Women of childbearing age?
  – Children and adolescents?
  – Persons living with HIV?
Activity 5: Role play: Management

Activity 5: Role play

- A health-care provider assessed this person and their spouse and decided that the person has epilepsy.
- The health-care provider now has the responsibility of developing a treatment plan with the person.
- The treatment plan should include psychosocial and pharmacological interventions as well as instructions to the spouse on how to help the person if they have a convulsive seizure at home and when to refer for medical help.

See EPI supporting material role play 2.

Print off the four different instruction sheets for the participants playing the different roles.

Ensure the person playing the role of the observer also has a competency assessment form (see ToHP training forms) in order to assess the participants.

Duration: 40 minutes.

Purpose: To enable participants to practise using recommended psychosocial and pharmacological interventions for epilepsy.

Situation:
- A health-care provider assessed this person and their spouse and decided that the person has epilepsy.
- The health-care provider now has the responsibility to develop a treatment plan with the person.
- The treatment plan should include psychosocial and pharmacological interventions as well as instructions to the spouse on how to help the person if they have a convulsive seizure at home and when to refer for medical help.

Instructions:
- Divide the participants into groups of four.
- Instruct one person to play the role of the health-care provider, one the person seeking help, one the spouse and one the observer.
- Distribute the role play instructions to each person depending on their role.
- Ensure that the participants keep to the allotted time.
Highlight the recommendations on frequency of contact (page 67 mhGAP-IG Version 2.0) and explain that follow-up should occur every three to six months.

Ask participants why they think that is?

Talk through step 1 of the follow-up algorithm and ask participants to brainstorm what questions they could ask at follow-up?

Possible questions could include:
- Has the person been keeping a seizure diary?
- Have there been any drug specific side-effects?
- Are they taking their medication as prescribed? If not, why not?
- Are they having any other issues?

Describe what to do if the person is not improving on their current dose, highlighting when they should refer.

Describe what should be monitored at regular follow-ups.

Focus on asking about how well they are being accepted and treated by the community.

Describe when to consider stopping medication and why.
Activity 6: Group discussion: How to reduce stigma and discrimination

<table>
<thead>
<tr>
<th>How to reduce stigma and discrimination?</th>
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<tbody>
<tr>
<td>1. Why is it important that you respect, protect and promote the rights of people with epilepsy?</td>
</tr>
<tr>
<td>2. Can you think of some concrete actions that you could undertake to make the rights of people with epilepsy a reality?</td>
</tr>
<tr>
<td>3. What would be the positive impact of these actions for all the groups concerned?</td>
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</table>

**Duration:** 30 minutes.

**Purpose:** To have participants reflect and plan what they can do to help reduce stigma and discrimination against a person with epilepsy and their carer.

**Instructions:**
- Divide the participants into three groups.
  - One group will represent people with epilepsy.
  - One group will represent non-specialized health-care providers.
  - One group will represent the family and carers of people with epilepsy.
- Give each group three pieces of flip chart paper and pens.
- You are going to ask the groups three different questions.
- They should write down their answers to the questions on three separate pieces of flip chart paper.
- Instruct the participants to write down their answers imagining that they are a person from the group they represent.

**Question 1:** Why is it important that you respect, protect and promote the rights of people with epilepsy?

**Question 2:** Can you think of some concrete actions that you could undertake to make the rights of people with epilepsy a reality?

**Question 3:** What would be the positive impact of these actions for all the groups concerned?
Possible answers to look for:

**Question 1: Why is it important that you respect, protect and promote the rights of people with epilepsy?**

**Potential answers from people with epilepsy:**
- We can contribute a wide array of expertise, skills and talents and these can benefit everyone.
- We are human beings and should have the same opportunities as everyone else.
- We know what is best for us; what is helpful and what is not helpful.
- We have the right to participate in all actions and issues that affect us.

**Potential answers from health-care providers:**
- I want to give the people under my care the respect they deserve.
- It is my legal obligation.
- This is part of my job and responsibility.
- It is the right thing to do.
- By providing care and support that respects people’s rights, people are more likely to accept the service we provide, respond well to our care and support and to recover.

**Potential answers from family members and carers:**
- I can help voice the wishes and preferences of my relative and help explain these to others when needed.
- I want what is best for my relative and these rights give them the best opportunities to live a good life.
- I can have an important role in enabling my relative to live a more fulfilling life by respecting their rights, being more accepting and changing some of my own actions.

**Question 2: Can you think of some concrete actions that you could undertake to make the rights of people with epilepsy a reality?**

**Potential answers from people with epilepsy:**
- I can learn my rights and understand them.
- I can explain my rights to my peers, family and the community members.
- I can speak with local officials about the need to change.
- I can help other people in the same situation as me.
- I can talk about my experience to raise awareness about disability and human rights.

**Potential answers from health-care providers:**
- I can make sure that my clinical practice promotes respect and dignity and the rights of people with epilepsy.
- I can train and inform other staff about human rights and make sure that my colleagues also promote respect and dignity.
- I can talk to people about epilepsy in my work place so that they understand.
- I can speak to service management about taking action to improve treatment for people with epilepsy.
- I can speak with local officials about the need to change.
- I can make sure that people with epilepsy are involved and participate in decisions concerning running services for them.

**Potential answers from family members and carers:**
- I can explain their rights to my relative.
- I can make my relative feel that I respect them.
- I can try not to over protect my relative.
- I can make sure I listen and respect their views and decisions.
- I can support and encourage my relative to make decisions and become independent.
• I can make sure other family members/community members respect the rights of my relative.
• I can speak with local officials about the need for change and for the creation of the services that meet the needs of my relative and other people with epilepsy.
• I can raise awareness in the community to break down stigma, stereotypes and prejudices.

Question 3: What would be the positive impact of these actions for all the groups concerned?

Potential benefits for people with epilepsy:
• I would have greater independence and be less dependent on my family, friends and health-care provider.
• I would feel more empowered to take control of my own life and recovery.
• I would feel stronger.
• I would be able to develop new skills.
• I would be able to contribute my skills and talents to society and be more included.

Potential benefits for health-care providers:
• I would be able to provide better quality of care for individuals.
• I would see better outcomes for people so I would feel happier in my job.
• I would be able to improve services provided.
• The people to whom I provide care and support would be empowered.
• Relapse and dependency would be reduced.
• I can make the service a better place to work.
• People to whom I provide care would lead more fulfilling and independent lives.

Potential benefits for family members and carers:
• I would feel better and happier because my relative was better and had a better quality of life.
• I would have more time to pursue my own goals as I would need to spend less time caring for my relative.
• I would feel empowered to be able to support my relative and start breaking down prejudice and stereotypes.
• I would feel less stressed and have better mental well-being.
• I would feel empowered to be able to talk to local community leaders and decision-makers about respecting the rights of people with epilepsy.
• I would have a happier family as my relative would be able to engage more in family life.
Session 5.
Review

 соверш
 15 minutes

**Duration:** Minimum 15 minutes (depends on participants’ questions).

**Purpose:** To review the knowledge and skills gained during this training session by delivering MCQs and facilitating a discussion.

**Instructions:**
- Administer the MCQs (see EPI supporting material) to participants.
- Discuss the answers as a group.
- Facilitate a brief discussion answering any queries or concerns the participants may have.
EPI PowerPoint slide presentation

PowerPoint slide presentation available online at:
http://www.who.int/mental_health/mhgap/epi_slides.pdf

EPI supporting material

- Person stories
- Role plays
- Multiple choice questions
- Video links

Activity 3: mhGAP EPI module – assessment
https://www.youtube.com/watch?v=RUIRg555xl0&index=6&list=PLU4ieskOli8GicaEnDwe5Q6-yaGxhes5v

Supporting material available online at:
www.who.int/mental_health/mhgap/epi_supporting_material.pdf