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University College Cork, Ireland Coláiste na hOllscoile Corcaigh

# Ollscoil na hÉireann, Corcaigh National University of Ireland, Cork



University College Cork, Ireland Coláiste na hOllscoile Corcaigh

Living with Motor Neurone Disease (MND)

and Dysphagia – the personal experiences of people

with MND and their caregivers

Volume 2 (2)

Appendices

and

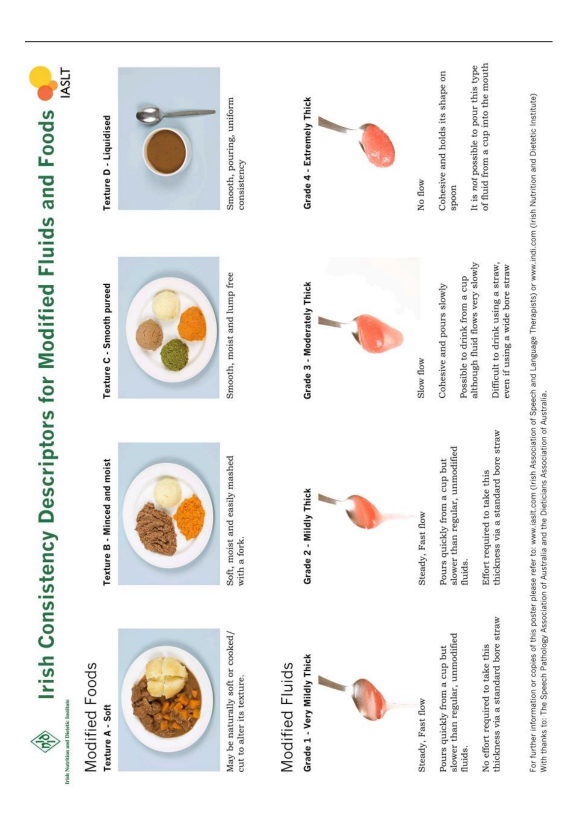
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Thesis presented by

Dominika Lisiecka, MSc. in Speech & Language Therapy

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### Appendix 1: Irish Consistency Descriptors for Modified Fluids and Foods

#### **Appendix 2: Functional Oral Intake Scale**

(Crary, Mann, & Groher, 2005)

### TUBE DEPENDENT (levels 1-3)

- 1. No oral intake
- 2. Tube dependent with minimal/ inconsistent oral intake
- 3. Tube supplements with consistent oral intake

### TOTAL ORAL INTAKE (levels 4-7)

- 4. Total oral intake of a single consistency
- 5. Total oral intake of multiple consistencies requiring special preparation.
- 6. Total oral intake with no special preparation, but must avoid specific foods or liquids items.
- 7. Total oral intake with no restrictions.

#### Appendix 3: ALS Severity Scale: Speech

(Hillel et al., 1989)

#### NORMAL SPEECH PROCESSES

**10 – Normal Speech**: Patients denies any difficulty speaking. Examination demonstrates no abnormality.

**9** – **Nominal Speech Abnormality**: Only the patient or spouse notices that speech has changed. Maintains normal rate and volume.

#### DETECTABLE SPEECH DISTURBANCE

**8** – **Perceived Speech Changes**: Speech changes are noted by others, especially during fatigue or stress. Rate of speech reminds essentially normal.

7 – **Obvious speech abnormalities:** Speech is consistently impaired. Affected are rate, articulation, and resonance. Remains easily understood.

#### **BEHAVIOURAL MODIFICATIONS**

6 – Repeats Messages on Occasion: Rate is much slower. Repeats specific words in adverse listening situation. Does not limit complexity or length of message.

5 – Frequent Repeating Required: Speech is slow and labored.Extensive repetition or a "translation" is commonly needed.Patient probably limits the complexity or length of messages.

#### USE OF AUGMENTATIVE COMMUNICATION

**4 – Speech Plus Augmentative Communication:** Speech is used in response to questions. Intelligibility problems need to be resolved by writing or a spokesperson.

3 – Limits speech to One-Word Response: Vocalizes one word response beyond yes/no; otherwise writes or uses a spokesperson. Initiates communication nonvocally.

#### LOSS OF USEFULL SPEECH

**2** – **Vocalizes for Emotional Expression**: Uses vocal inflection to express emotion, affirmation, and negation.

**1** – **Nonvocal:** Vocalization is effortful, limited in duration, and rarely attempted. May vocalize for crying or pain.

#### Appendix 4: Letter of Invitation to Participate in a Research Project

# Living with Motor Neuron Disease (MND) and Dysphagia – the personal experiences of individuals with MND and their caregivers.

My name is Dominika Lisiecka. I am a Speech & Language Therapist and a student in the Department of Speech & Hearing Sciences at University College Cork. I am conducting a research study as part of the requirements of my PhD degree. I would like to invite you to participate in this study. Please read this letter as you have to be fully aware of time commitment and any potential risks and benefits to make an informed decision.

I would like to study the experiences of people living with motor neuron disease (MND), who also have swallowing difficulties. I am contacting you as you have been diagnosed with MND and swallowing problems or you are caring for someone with these problems.

If you decide to participate, I will meet you to get your consent and I will check if you fulfil all criteria to take part in this study. After that, you will be asked to meet me for two interviews. There will be an option of scheduling more interviews if we both feel the need to do so.

The interviews will take place in your home. However, if you prefer to meet me at different location, I can provide an alternative venue.

The time will be mutually agreed and the duration of each interview will be 45 minutes maximum (for a person with MND) and 60 minutes (for a caregiver).

Each session will be audio recorded so that I can accurately reflect on what is discussed. If you use alternative communication I will ask for your permission to video tape the session, but you have right to refuse (and we will audio record only). The recordings will only be reviewed by members of the research team, who will transcribe and analyse them.

Your participation is confidential. Study information will be kept in a secure location at the University College Cork. The results of the study may be published or presented at professional meetings, but your identity will not be revealed.

You may feel emotional during the interview and you can stop the interview at any point. We will also stop when you get tired.

Taking part in this study is your decision. Although you won't benefit directly from participating in the study, we hope that others affected by MND and swallowing difficulties will benefit, as this project should improve our understanding of these problems. The data you will provide will be analysed, the result will be presented in a thesis and shared with relevant professionals. It is hoped that this information will result in improved quality of life and quality of care for both patients with MND and their families.

Your participation is voluntary and you can withdraw at any stage with no consequences. You may also request to have your data destroyed up to two weeks after the last interview.

There is no financial reimbursement for participation in this study.

I will contact you next week to see if you would like to participate. Alternatively, you may contact me at any stage and I will be happy to answer any questions. My phone number is 0861564494 and my email is MNDresearchPhD@gmail.com.

You may also wish to contact my Chief-Investigator, Dr. Helen Kelly at the Department of Speech and Hearing Sciences, School of Clinical Therapies, Brookfield Health Sciences Complex, University College Cork, College Road Cork, Ireland; tel: (0) 21 490 1746; fax: (0) 21 490 1542; email: helen.kelly@ucc.ie

With kind regards,

Dominika Lisiecka

#### **Appendix 5: Consent for Participation in Research**

#### **Information Sheet**

## "Living with Motor Neuron Disease (MND) and Dysphagia – the personal experiences of individuals with MND and their caregivers.

You are being asked to participate in a research study investigating the experiences of swallowing difficulties in Motor Neurone Disease (MND). In order to decide whether or not you want to be a part of this research study, you should understand enough about its risks and benefits to make an informed judgment. This process is known as informed consent. This consent form gives detailed information about the research study, which will be discussed with you. Once you understand the study, you will be asked to sign this form if you wish to participate.

#### **STUDY INFORMATION:**

We are doing this study to improve our understanding of swallowing problems in Motor Neurone Disease. We are interested in finding out how people living with MND experience swallowing difficulties. We would like to talk to adults diagnosed with MND and their caregivers.

#### Interviews

At least two interviews are planned with every person. The duration of each interview will be approximately 45 minutes for the person with MND and 60 minutes for the caregiver. We will voice record all interviews. In some circumstances we may ask you to make a video recording, but you have the right to limit the recording to voice recording only.

#### **Inclusion criteria**

We have a list of specific conditions, which you will need to fulfil in order to take part in this project. For example, we will look at the time since your diagnosis, what diet you eat/ prepare, what your thinking processes are like. We will explain all conditions to you at our first meeting. If you fulfil the conditions we will schedule our first interview.

#### What will happen to your records?

Ensuring that your records are kept confidential is very important to us. We will remove your name and any other information which could identify you. Only the research team may access the original data. The recordings of our talks will be stored on an encrypted laptop / encrypted external drive and kept in a locked filing cabinet in a locked room at the University College Cork.

We will analyse your recordings and results will be presented in a thesis and shared with relevant professionals. Some of the interview (for example, quotations) may be presented at conferences or published in writing, but your identity will never be revealed.

#### Is the information I share in the interviews confidential?

The information you share in the interviews are considered confidential. However, in some exceptional circumstances it may become necessary to break confidentiality. Exceptional circumstances include:

1. If there is a serious concern that there may be a threat to the safety or life of you or others.

2. In the context of criminal behaviour and disclosures required by legal process.

3. In the context of child protection and elderly protection issues.

If this is required, it will be done with sensitivity and on a need to know basis. Your interviewer will make every effort to discuss this with you before that step is taken.

#### Potential risks and benefits

You may feel emotional or tired during the interview and you can stop the interview at any point.

Taking part in this study is your decision. Your participation is voluntary and you can withdraw at any stage with no consequences. Your withdrawal will not have any impact on the health services normally available to you. You may also request to have your data destroyed up to two weeks after the last interview.

Although you may not benefit directly from participating in the study, this project may improve our understanding of these problems. It is hoped that this information will result in improved quality of life and quality of care for both patients with MND and their families.

#### **Primary Investigator:**

#### Dominika Lisiecka

Department of Speech and Hearing Sciences, School of Clinical Therapies, Brookfield Health Sciences Complex, University College Cork, College Road Cork, Ireland

Tel: 0861564494

Email: MNDresearchPhD@gmail.com.

Dominika Lisiecka is carrying out this project as part of the requirements for her PhD at University College Cork which has been funded by the Health Research Board. She is being supervised by Dr. Helen Kelly and Prof Jeanne Jackson.

#### **Chief-Investigators:**

Dr. Helen Kelly, Department of Speech and Hearing Sciences, School of Clinical Therapies, Brookfield Health Sciences Complex, University College Cork, College Road Cork, Ireland Tel: (0) 21 490 1746 Email: helen.kelly@ucc.ie Prof. Jeanne Jackson, Department of Occupational Sciences and Occupational Therapy, School of Clinical Therapies, Brookfield Health Sciences Complex, University College Cork, College Road Cork, Ireland Tel: (0) 21 4901536 Email: j.jackson@ucc.ie

#### AGREEMENT TO CONSENT

1. I confirm that I have read and understand the information sheet for the above study and have had the opportunity to ask questions.

2. I understand that my participation is voluntary and that I am free to withdraw at any time, without giving reason.

3. I agree to take part in the above study

4. I agree to the interview being audio recorded

5. I agree to the interview being video recorded (if appropriate)

6. I agree to the use of anonymised quotes in publications

7. I understand that in exceptional circumstances where a significant risk of safety of life has been identified it may be necessary to break the confidentiality. This will be discussed with me before any action.

I have received a copy of this consent form for my records. I understand that if I have any questions concerning this research, I can contact the persons listed above.

After reading the entire consent form, if you have no further questions about the study and wish to participate, please sign where indicated.

I, the undersigned, hereby consent to participate in the above described project conducted at University College Cork.

Signature of participant:\_\_\_\_\_

Date and time: \_\_\_\_\_

Witness:

Date and time:

#### Appendix 6 a): Interview Template for a Person with MND

Name: \_\_\_\_\_

The following background / demographic questions will be asked following the interview in an open-ended manner (as far as possible)

DOB:
Address:
GP:
Neurologist:
Medications:
Member of IMNDA: Yes  No
Current diet: Oral $\square$ Non-oral $\square$ (NG $\square$ PEG $\square$ PEJ $\square$ REG $\square$ ) both $\square$
Current liquid consistency:
Current food texture:
Self-feeding: Yes □ No □
Occupation:
Lives with:

Interviewer introduction:

Hi, my name is Dominika and I am a student at UCC. As part of my PhD I am conducting a research project and as part of that I will be interviewing you today. I am very glad that you gave your consent to participate in my project. You can stop the interview at any point. As discussed before, I will record your voice to help me analyse our interview. Please let me know if you feel tired and need to take a break.

Main questions	Additional questions
I am interested in getting information about you and your life. Can you tell me a little bit about you and your family?	Do you work? (if appropriate) How is work?
If I followed you throughout a typical day, what would I see you doing?	
Some people say that they have good days, days when they have good energy and are able to do more, and bad days, days when the energy is lower and they are not as active. What about you?	
We are going to talk about food and eating now.	
Can you tell me about your typical breakfast, lunch and dinner?	Can you elaborate on that? That's helpful. I'd appreciate more details.
I would like you to go back to the past. Can you recall any meal that you really enjoyed? What was it like?	When was it? Where were you? Who were you with? How did the food taste/ smell/ look?
Some people love food and food is very important in their lives. Others eat to keep going, but don't really pay much attention to food. What about you? What place does food have in your life?	
In the last few months has anything changed in the way you eat or drink? Can you tell me more about this?	
Can you tell me how did you discover that your swallow had gotten weaker?	Alternative: How has the way you eat and drink changed in the last few months?
What knowledge did you have about the swallowing before you became sick?	
How do you understand swallowing problems now?	
What is it like to have swallowing problems and MND?	Suppose I was someone who never heard of swallowing problems and MND, how would you explain these to me?
If you had any questions related to your swallowing, who would you ask?	
What do you think about the services you received for your swallowing problem?	
If you had a power to make any changes to the services you and your family received so far, would you change anything?	(If yes) What would you like to change?
Conclusion of interview	
Would you like to add anything else? Were you surprised by the questions I asked? Did I leave anything out that you thought I would ask?	

#### Appendix 6 b): Interview Template for a Caregiver of a Person with

#### MND

Name:	_
Date:	
Address:	
Member of IMNDA: Yes $\square$ No $\square$	
Occupation:	
Relationship with the person with MND	

Introduction of interviewer:

Hi, my name is Dominika and I am a student at UCC. As part of my PhD I am conducting a research project and as part of that I will be interviewing you today. I am very glad that you gave me your consent to participate in this project. You can stop the interview at any point. As discussed before, I will record your voice to help me analyse our interview. Please let me know if you wish to take a break.

Main questions	Additional questions
I am interested in getting information about you and your life. Can	Do you work?
you tell me a little bit about you and your family?	How is work?
If I followed you throughout a typical day, what would I see you doing?	
I would like to talk about X (person with MND) now.	
Can you tell me how you care for X?	
What is it like to look after X?	
Suppose I was someone who is just about to start caring for someone with MND. How would you help me to prepare for this role?	
When you care for X, is there anything you find particularly hard?	(if yes) Is there anything that you find the hardest?
I would like to talk about eating now.	
Can you describe X's typical breakfast, lunch and dinner.	
How exactly do you help X with eating and drinking?	How do you prepare food for X?
I understand that X has some swallowing problems. Can you tell me how it all started?	Can you take me through what happened?
In the last few months, has anything changed in the way X eats and drinks? Can you tell me more about that?	Alternative: How exactly has the way X eats and drinks changed in the last few months?
What is it like to look after someone with swallowing difficulties?	
What knowledge did you have about the swallow before X became sick?	
How do you understand swallowing problems now?	
If you had any questions related to X swallowing, who would you ask?	
What do you think about the services you received for X's swallowing problem?	
If you had a power to make any changes to the services you and X received so far, would you change anything?	(If yes) What would you like to change?
Conclusion of interview	
Would you like to add anything else? Were you surprised by the questions I asked? Did I leave anything out that you thought I would ask?	

Superordinate theme	Subordinate themes	Emergent themes	Quotations	Line
1.	1.1	"I thought	What did you feel? [when the	16-17
Living with	Pre-	I was going to die"	swallow problems started]	
dysphagia	gastrostomy		I thought I was going to die.	
	experiences			
		Swallow	So Sally, can you remember when	12-14
		deterioration	the swallow problems started?	
			Ehm (pause) I don't know. About 4	
			weeks before the tube.	
		Losing	() all my clothes got too big.	115-
		weight		116
		Choking	And can you remember the first time	18-26
			when the choking happened to you?	
			I can.	
			What was happening? Can you	
			describe?	
			I couldn't get my breath.	
			What were you eating?	
			I can't remember.	
			Were you on your own?	
			Maybe? It was nothing. Got worse.	
		Unable to eat	And were you advised not to take	84-85
			anything by mouth?	
			I can'tcouldn't (body language	
			indicating no ability to swallow at all)	
		Determined to look	What were you thinking before? Did	31-32
		for explanations	you have anything in your mind	
			what was happening to you?	
			I was thinking we have the money to	
			go anywhere in the world to get help.	
		No	And what knowledge did you have	95-96
		previous knowledge	about the swallow before?	
		of dysphagia	No.	
		Diagnosis	Can you remember your first visit to	27-30
		as a relief	[hospital]?	
			Yes	
			How was that for you?	
			It was such a relief to know at last what	
			was wrong with me.	

#### Appendix 7 a): Data Analysis Table: A PwMND (Sally)

#### Comments:

This was a very stressful time. The time of not knowing what was happening with her body. Her speech started to slur and she became unable to swallow. She was not able to eat anything at all for about 2 weeks before she was diagnosed. She thought she was dying, but she didn't give up. She wanted to find out what was wrong with her. She was prepared to travel and pay a lot of money to get some answers. There were delays between visits with consultants. Finally, Sally's daughter rang the IMNDA and she got an appointment within two weeks. Sally went up to the MND clinic and got her answer: MND. Not many people would say that they felt relieved when they heard the diagnosis. For Sally, the diagnosis explained

her symptoms and gave her access to medical management and support. Maybe she felt understood and reassured.

Sally has a good insight into her swallowing status. She is not denying the problem. She admits it. She remembers her first choking experiences but that was only the beginning. The swallow deteriorated with time to the point where Sally became unable to swallow anything. Even a taste of food was not possible. Her own saliva became a serious risk and could potentially choke her.

Would it have helped Sally to have had some previous knowledge of dysphagia? Would this have helped her to find the answers quicker? It must have been so exhausting and hard. She continued her house-hold routines, such as shopping and cooking and she was trying to eat but it was becoming harder and harder. She did not suspect MND. In fact, she did not suspect anything. She had no clue. She couldn't understand what was happening to her and she was convinced that she was dying.

When I think about Sally's life before she got the diagnosis and the gastrostomy tube (which was almost simultaneous), I feel that she escaped death in the last moment. Prior to non-oral feeding she was already unable to eat and lost much weight. Paradoxically, as how can you say that someone has escaped death by being diagnosed with a terminal disease... but this was the case for Sally. In her circumstances, the diagnosis of MND opened the door to the medical services that she required. Although Sally could not be cured, her symptoms could be managed, ultimately prolonging her life.

	1.2	Acceptance of	And was it [doctor] who talked to	38-43
<mark>.</mark>	Living with	gastrostomy	you about the tube?	
	non-oral		Hmm [yes]	
<mark>1</mark>	feeding		Can you remember that	
			moment?	
			[nod]	
			What did you think of it?	
			I was willing to do it.	
		Non-oral	What was your impression when	50-53
		feeding as a	you saw the tube for the first	
		relief	time?	
			[vocalisations]	
			Is that nervous?	
			[Vocalisation]. [wrote]: Relived.	
		Feeding regime	We are working on it now.	113
			And I understand the feeding	67-70
			takes a long time now.	
			Hmm (yes). 16 hrs.	
		Position	Liam cleans the RIG and I come out	68-69
			and sit for the rest of the food to	
			finish.	
			I would like to be able to sleep on	117-
			my side and put my arm around	121
			Liam.	
			Is that not possible because	
			The pillow is very high and I have	
			to sit up.	
			Has that changed the way you	
			sleep?	
			Yes.	
		Husband's	If I followed you through you	65-68
		support	typical day what would I see you	
			doing from the moment you wake	
			up?	

	Liam takes out the food thing and I	
	have a shower	
	Liam cleans the RIG ()	
Problems	So what happened today? You	4-5
	had to go to the doctor	
	I broke the tube.	
	My tube came loose.	

Sally was willing to be fed non-orally. She was willing to live and she had no other option. Her swallow had failed completely. It was a life and death choice for her. Her initial reaction when she saw the RIG for the first time was the feeling of being relived. The procedure was successful. The gastrostomy tube was there. In her stomach. A small piece of plastic which could prolong her life. She had just managed to get it on time. This was due to her daughter who contacted the IMNDA. Although Sally had been seen by various medical professionals the progression of MND was so much faster than the delays in the health service. Sally tried to get help. She was told to wait. Each time. Her symptoms were classic and although the diagnosis could not have been confirmed without undergoing specific tests, the doctors must have had some suspicions. How was she left starving for 2 weeks before seeing a doctor, who decided to perform a gastrostomy, (as reported by her husband)?Sally had been seen by many professionals for many weeks, yet when her dysphagia progressed dramatically leaving Sally unable to eat and drink, no one was around to help her.

Although the feeding tube prolonged Sally's life, it came with specific problems and demands. The feeding regime took time to establish. It was a trial and error approach where various types of feeds were given to see how Sally's digestive system reacted. She still requires to be fed very slowly. Although she is being fed all night it is not sufficient. Her feeds take 16 hours to be administered. Sally needs her husband's support to look after the "food thing". She has lost a lot of independence.

Re: position. It is known that non-oral feeding increases the risk of reflux and subsequently chest infections or pneumonia. Having to be fed overnight requires a person to be in an upright position to minimise the occurrence of reflux. This is the medical perspective. For Sally, having to be upright at night affected her sleep in many ways. It prevented her from being able to "put her arms around" her husband. She missed that. Sally was medically safe at night but emotionally unsafe.

Professionals often consider and recommend sleeping position for patients. Should they look beyond the medical reasons and consider how an altered sleeping position affects the person holistically?

Problems with the gastrostomy tube: When I arrived one morning Sally was not at home despite asking me to call to her. There was an emergency – the tube became loose and they had to rush to the doctor. Sally was hoovering and maybe bending caused the tube to dislodge a bit. When she said "I broke the tube" I thought she felt guilty about it. She knows that the tube extended her life and maybe she felt that she should be more careful with it. She is confined to the house due to the long feeding time and she wants to stay active. She attempts to participate in her house chores, but the tube is there to stop her a bit. It is there to remind her to slow down and be mindful of herself. The tube is a gift of life and wants to be respected.

remina ner to stow down and be minaje	ai of herself. The the	re is a gift of life and wants to be respec	icu.
	Social impact	We used to go for meals out and	59-60
		have friends in but never I cannot	
		do that.	
	Missing food	I would love to have a rasher	82
		sandwich but I cannot take any	
		by mouth.	
	Food as a	And in the past would you say	89-92
	means to	the food was important in your	
	socialise	life?	
		Yes.	

	How important was it?	
	Well it was always good to	
	socialise.	
Changed	I was 80 last week. All the children	125-
participation in	come and we were all together.	127
celebrations	Lidia had a nice meal just the	
	but I could not eat.	
	Or have a glass of champagne.	

Sally's life used to evolve around food. She was a lively housewife who cooked a lot and invited people for meals "while Liam was making the money". That's how they shared their duties: Liam was working outside the house and provided finance for the family, Sally looked after the house and family's social life. Food was important to her mainly as a way of socialising. Now she has lost her ability to swallow and she has lost her social life.

In our culture celebrations involve food. The food becomes symbolic: we expect a cake for someone's birthday or wedding, we cook turkey for Christmas, and buy chocolate eggs for Easter. We celebrate occasions with champagne. It becomes so difficult when a person is unable to swallow. What to do? Although Sally is being fed non-orally, other family members continue to eat in the natural way. When they meet together to cerebrate various occasions, there really is no way to avoid eating and drinking. What can be a substitute for Sally who is unable to even taste the food or drink? Her daughter made a meal for Sally's birthday, but it was not for Sally. They celebrated Sally's 80s with a glass of champagne, but she could not participate. What could have been done to help Sally celebrate her birthday and to have something special that day? What would she like?

The family will meet in Sally's house this year as she feels unable to go to her daughter's house as before. This is because of non-oral feeding. It is still new to them and Sally does not use it outside her house. Sally will cook Christmas dinner. I am sure she has everything well planned. When she said "some dinner" it sounded like she is not so excited about the food anymore, she cannot be. It is not for her. She will make an effort to cook well for others, because she loves them. There is nothing she can make for herself, nothing that she could try at the Christmas table. Maybe she had engaged in the process of detaching from food. It was never going to be a part of her life again.

When you have severe dysphagia you are not able to participate in the celebrations as before. It is a fact. We can look for substitutes but it will not be as before.

1.3	Lost	Lost skills	I would not be able to do all myself.	115-
ind	ependence		The not being able to talk. Or go	116
			out myself	
		Requiring	So how has your life changed now	59-60
		constant support	since the diagnosis?	
			Well I cannot even do shopping	
			without Liam. I have to have Liam	
			all the time.	
		Being a burden	I have to have Liam all the time and	60
			he gets very tired because of me.	

Comments:

Sally lost her independence despite being fully mobile and having fully preserved limb functions. She has no speech and no swallow and she knows that the MND can cause further deteriorations at any time. She does not go out by herself due to her inability to communicate and she needs a lot of help with non-oral feeding

at home. It shows how people' circumstances differ. Other participants with bulbar onset of MND reported that they were still very independent. Sally cleans and cooks at home. She can dress herself and shower. She is independent in self-care except for non-oral feeding. I am not sure why she has not attempted to manage it herself. From the beginning it was Liam who looked after it. He was trained in the hospital and he continues to manage non-oral feeding, although Sally's fine motor skills are perfect.

Sally is living a different life with MND. No food or drink. No social life. Participation in family cerebrations/ occasions are not the same. No independence. So vulnerable. Unable to hug her husband at night. Unable to chat to him. Maybe her self-worth suffered too? She says that buying new clothes would be a waste. Is it that she is not worth them as she is not leaving the house or having many visitors, or is it that she feels her life can come to an end soon?

1.4.	Empathy	Ann [therapist] was very nice but	37
Professional	I ···· J	Jane [another therapist] was more	
attitude		understanding	
1.5 Making	Approaching	and I think if I bought new clothes it	115
sense of own	death	would be a waste ()	
suffering			
	Appreciation of life pre-MND	We have had a good life and I was going to offer any pain or discomfort to God in thanksgiving for all.	54-55
	Suffering as thanksgiving	We have had a good life and I was going to offer any pain or discomfort to God in thanksgiving for all.	54-55
	Offering discomfort to God	We have had a good life and I was going to offer any pain or discomfort to God in thanksgiving for all.	54-55
	Importance of religion	Okay. How important is religion for you? Very.	56-57

#### Comments:

Sally's life is very hard for her. It changed a lot. She lost a lot. However, Sally decided to do the best she can. She continued being involved in her house duties including cleaning and cooking. She consented for non-oral feeding; she was not ready to die. Sally contrasted her life with MND to the "good life" before. She feels grateful for her past and she offers her current suffering to God as a thanksgiving for what she had got/ experienced before. This altruistic approach helps her to cope with MND. It added sense to her suffering. It made it easier to accept it. Religion helped her to focus on "doing the best she could" instead of thinking about ending her life and giving up. "The not being able to talk" is particularly hard for her. Sally uses a lot of body language to communicate.

Sally is so busy at home, she moves so fast. She used to exercise and she is still very fit. Her baking and cooking is perfect. There is no hesitation, no mess. Everything seems to be well planned. Years of experience. But when I look into her eyes it feels like the spark is gone from her. There is no voice and there is no independence. She is doing her best given her circumstances, but I don't feel she is very happy. She is trying to use her time well, but she feels insecure. She is also aware that her husband gets tired looking after her, she is a burden for him now. He is older and she used to care for him all their lives. I don't think she regrets her decision to commence non-oral feeding as she was not ready to die, especially to die of

something unknown and unexplained. She wanted to continue living and she accepts the consequences. There is a price you pay for everything in life. She is pragmatic and realistic. In the past, Sally was able to stay at home with her children as the husband provided for the family. She travelled around the world. She had hobbies. She raised four children and is very proud of them. She has her grandchildren and she loves them all. Towards the end of her life she says THANK YOU by accepting her suffering. Her memories are still alive and no one can take them away from her. They help her to keep going. The religion adds sense to her changed life and it provides some sense of peace when thinking about the end.

After this interview was completed I posted Sally a notebook with follow-up questions:

- 1. How have you been since I saw you? How is Liam?
- 2. How did you spend Christmas?
- *3. How is the RIG going?*
- 4. How does your swallowing feel now?
- 5. What do you think about the services you received for your swallowing?
- 6. Is there anything else you would like to add?

When I did not receive the notebook by the suggested deadline I presumed she was unable/ not willing to continue (there had been changes in the family's circumstances which prevented me from calling Sally's relatives to find out about her willingness to complete the study).

On the 13<sup>th</sup> of April (2016) I received a phone call from Sally's husband. He apologised for not sending the notebook back to me. Sally had completed it within the specified deadline, but he misplaced it. We scheduled another interview, as per Sally's wishes.

There was almost a 5 months gap between the interviews and (after consulting my Supervisory Team) I therefore decided not to include the 2<sup>nd</sup> in the further stages of this study. I validated the findings from my first interview. Sally reported that she got used to not eating and it "does not bother her" to see food anymore (so she does not desire to have her bacon sandwich anymore). I also perceived a change in her relationship with Liam. I feel he is becoming increasingly less able to look after Sally due to his own health issues. Sally appeared worried that he may die first and what would happen to her. She also provided information about Liam becoming less tolerant of her disabilities, for example, he continuously asks her to close her mouth while she is not even aware that her mouth is open.

She also said that the sleeping is going well and there was no mention of wanting to hug Liam at night anymore. So there were changes in the emerging themes which could change the overall interpretation of her experiences. However, the information pertaining to perceiving the diagnosis as a relief, offering suffering to God, and lost independence was still confirmed.

After the first interview my personal overall theme describing Sally's life was: the experience of love in MND. After the second interview this theme would change to: how progression of MND killed love.

## Appendix 7 b): Data Analysis Table: A Caregiver (Jackie)

Superordinate themes	Subordinate themes	Emergent themes	Quotations	Line
1.         Desire to         prolong her         husband's life	themes1.1.Supportingher husbandall the time	Prioritising Chris's needs	That's what we did, you know, we make the most of every day and make sure that anything Chris wants to do he can do it. We went to a healing mass last night now. Yeah it was kind of a healing ceremony. My brother rang and said it was on so we dropped everything and we went, em, like Chris will want to go to the match this evening, so we will make sure that Chris goes to the match whoever else goes	653- 655 80-83
		"We're all running around him"	everywhere else. And usually we're racing (laughs), running around. Well Chris isn't. He spends all of his time sitting here now but we, em, we're all running around him so like going, here, there, doing everything but when you go now the match will have to be sorted	521- 523
		Determined to protect Chris	I saw a fella going for his pocket and he did (laughs) and I was very bold. I just pushed the crowd in front of me, the whole lot of them onto the tube because I could see this fella aiming at Chris. I just gave a big push and nobody was going to get injured like. I just pushed the crowd like, you know, because he was two ahead of me and I pushed the crowd and we got onto the train and he was giving out mad. Who was pushing us and I said nothing and then, eh, and we were all just on the train then and everybody was pushing and the door was just about to close and the fella who did try to steal the thing ran off like a bat. He definitely because I was saying did I imagine that? No I didn't. He was just gone.	595- 603
		Ready to give up work for Chris	I would give work up in the morning like if Chris wanted me and that is the situation. I said	173- 175

Chris when you want me to stay at	
home say so and I'll stay at home	

She may be delicate and softly spoken. She may be fragile and heart broken. However, when it comes to protecting her husband she gathers her powers and becomes unstoppable and fearless. She has a clear priority: to protect her family. Chris is vulnerable now due to MND so she does her best to mind him and to ensure his safety all the time. She wants him to live as long as possible. They are both young (early 50s) and they have (...) children. I feel that prolonging his life is her goal and she is doing her best to achieve it. She knows that Chris doesn't have much time left therefore his needs and wants are the priorities.

When I was in their house I could feel the strength of this family. The way they communicated with each other, the way they spoke with me. I met two of Jackie's (...) children. They were not only so kind and nice but also appeared to be strong, mature, and very independent.

	and very macpenaenti	
Ensuring	I might make the soup but I'd put a	218-
appropriate	lot of cream in Chris's, you know, to	221
diet	keep the calories because like I was	
	told to maintain his weight. On day	
	one the first thing, one of the nurses	
	she says just keep the weight on.	

Comments:

Jackie looks after Chris's diet as she knows that maintaining his weight means longer survival. She is also aware of his swallowing difficulties, therefore, she ensures that food consistencies are safe for Chris. She is modifying food to help him stay as healthy as possible, not because she was advised by a SLT. She has a clear understanding of the relationship between food and Chris's health status. Extra calories are required to maintain his weight (can you really battle this with muscle wasting?) and soft food means less choking. Her main motivation is to keep Chris alive and well as long as possible.

5 0			
	Looking after	I suppose like obviously Chris isn't	175-
	husband's	working at the moment and I	177
	mental	suppose the only reason we're	
	wellbeing	keeping on the cows at all and	
		working hard is to keep state of	
		mind good (upset)	

Comments:

Jackie knows her husband well and she respects and acknowledges his needs (also psychological). I think she is trying to keep Chris's spirit up to give him more motivation to live and keep adapting to what MND requires from him (non-oral feeding, mobility aids, losing independence, etc.). I think she hopes that by ensuring that Chris is as happy as possible she can motivate him to live longer (to want to live longer). She tries to continue their "normal" (previous) life style. Chris worked on [anonymised] all his life and she knows that selling [it]m would come as a huge shock to him. She took on Chris's duties and she gets some help from her children. This is to avoid this shock (the shock of selling the [anonymised]) for as long as they can. I feel that if Chris loses his motivation to live he may not consent to having a gastrostomy or any other support. She wants him to see the point in living, despite his progressive disabilities.

Spending time	I'm last in the door to work so I just	50-54
spending time	I III last III the door to work so I just	50-54
with her	stay at home until then and have	
husband as	breakfast with Chris and then go to	
much as	work at nine and, em, I work until	
possible	three o'clock and I come home	

straight at three as well. Again I used to stay in work until four	
So, I don't know. I was there [at	424-
dysphagia training] that day now	426
without Chris and I didn't feel any	
better. () I felt it was a waste of my	
time being away from Chris to be	
honest.	

Jackie's routines have changed. She is so focused on being with her husband as much as she can that everything else is secondary. She spends her time at home with Chris as much as possible. She goes to work at the "last minute" and she drives home straight after. She needs to have a good reason to go somewhere without him. She doesn't sleep as well or as long as before.

I feel her pressure – unstoppable, constant, and exhausting.

Jackie's holiday routine has changed too. They used to be so active when travelling. They used to go abroad annually in recent years. This is not possible anymore. The way they plan has changed for her. "It is more day by day". This must influence her perspective on life. She doesn't appear to have anything to look forward to. She is constantly worried and stressed. Even if they were to book a holiday now "for the sake of the children" Jackie would worry whether Chris would be able to go. MND is like an unwanted shadow. It is there all the time and it follows you, your movements. You may not be able to see it on a very bright day, but it is always there.

Although Chris has always been a good driver (and he still is according to her), Jackie noticed that people are becoming more nervous about him driving as they see his inability to talk. She realised that when "last day now one of the neighbours said she'd go with him because I couldn't go I had to milk the cows and she, em, she sat in with him to go somewhere and I'd say he doesn't know what to think now (...) He's was an excellent driver always and he still is. I suppose people begin to get nervous of him so I don't know what's happening". This is upsetting for Jackie. For her, people could potentially stop Chris from doing what he has been doing for a long time and what he loves doing. This must be so upsetting for him too. There are things he is unable to do anymore and it is hard enough to live like that. There are other things that he can still do, but society doesn't trust his skills anymore. This is so cruel, yet somehow I can understand it. The nature of MND is to blame here. People may know the disease will progress, but no one knows when. People may fear that Chris may became unwell when driving their kids to the match. Jackie feels confused ("I don't know what's happening") and she finds it hard to understand why people are nervous about Chris's driving. She is convinced his driving skills are perfect especially when it was confirmed by an OT. I think she will do her best to be in the car with Chris all the time so he is not being joined by neighbours who don't trust his skills. This would bring an additional burden for her as, for example, she would have to milk the cows at a later time or get someone else to do it (so having to pay for it while she is the only earner in the house now). She may do it as she is focused on Chris's needs and she want to support him to do what he still enjoys. This is all contributing to his willingness to live and you need that to try to adapt to MND

ity to adapt to MINE				
	1.2. Non-oral	PEG as a	What made him decide [for	355-
	feeding as life	"good thing"	gastrostomy] do you think?	363
	prolongation		Well, em, it was getting harder to eat	
			like. I suppose I did say to him Chris	
			we want you around as long as	
			possible because we had heard like	
			that it will prolong life, you know, at	
			this stage it will prolong life. Chris	
			what do you think and one day I	
			didn't really push it hard. I didn't	

Anxious for Chris to undergo gastrostomy	really push it and I didn't push it at all but I did say it will prolong life and its good for, em, to keep all your body strong and em, and I said we want you around as long as possible. ()And one day he said okay I'll go for the tube. But like I did ring and suppose I be ringing again today. I didn't ring now yesterday but it is on my mind. (Long pause). I thought really when he made the decision like because there was so much, people saying to us like you should get the tube, you should get the tube. I thought that when he decided that he would get the tube. And here we are now six weeks later and after coming to this monumental decision in our lives	444-450
	monumental decision in our lives we're still waiting.	
Aware that Chris may become too unwell for gastrostomy	And I'm also very aware of the fact that after a certain stage you can't get the tube like. So I actually, I don't know. I worry sometimes is he still alright to get it but I hope he is.	440- 442

Jackie sees the PEG tube as the opportunity of have her husband alive for a longer time. She believes "it will prolong life" and that the feeding tube is "a good thing". She really would like Chris to undergo a gastrostomy as she knows it may become necessary in the future. When I was leaving her house one day she told me about Chris's (...) who died of MND. "He really died of starvation" she said.

Jackie knows there is no cure for MND and there is very little that can be done for her husband. Possibly the only hopeful news she heard from the professionals was in relation to the gastrostomy tube. This may be the only / one of the few ways of taking some control over the progression of MND. This is not a cure, but it matters. It is a solution for her. She remembers that:

"So then eight weeks ago, Chris's uncle came down and he spoke quite forcibly about it and I said and then a dietician had said it to us as well and the speech therapist said it to us"...

Some professionals recommended the PEG for Chris and Jackie heard "it may prolong life". At the same time she wanted Chris to decide for himself as she doesn't feel it would be respectful and appropriate to influence his decision.

"One day coming down in the car then I said like the speech therapist took out the tube again to Chris and I said what did she say about the tube Chris and he said leave it all back there (laughs) and I said no more."

Jackie is also aware of specific conditions that Chris needs to fulfil to be medically safe for the tube insertion.

"So Easter, Holy Thursday then I rang again because Chris was finding it tough to eat and I was afraid the time would pass because there's only a certain amount of time, you know, you can get it so I rang again and I rang again last week and left a message again and then we went up yesterday".

She is anxious that it may be too late for Chris to get the tube. She was hoping to get some help from professionals to convenience Chris to go for the tube, but not everyone was helpful to her in this regard:

"I thought going up that day that, em, oh great now she'll [the doctor] tell him get the peg and all will be great and but anyway no she wasn't... She didn't push the peg at all".

According to her priority (willingness to have her husband alive as long as possible) Jackie changed her behaviour from "I said no more" to verbalising her opinions. There was no time to remain silent anymore. It was hard for her: feeling the urge to influence Chris's decision while strongly believing that the choice should be purely his. She noted the deterioration of Chris's swallow and she could not hide her feelings anymore:

"It was getting harder to eat like. I suppose I did say to him Chris we want you around as long as possible because we had heard like that it will prolong life, you know, at this stage it will prolong life". She saw the tube as a tool to support her main priority: having Chris alive for as long as possible".

Jackie is almost explaining her behaviour to me and trying to justify it:

"I didn't really push it hard. I didn't really push it and I didn't push it at all but I did say it will prolong life and its good for, em, to keep all your body strong and em, and I said we want you around as long as possible".

She is justifying that her words were not to push Chris to decide, but she really wanted him to get the tube and she attempted to influence his decision at the same time.

2. Internal	2.1.	No choice but	() I suppose you come to live	138-
conflict	Maintaining	to accept	with it a bit then. Well you just	139
	a positive	MND	have to accept it and try and go on.	
	attitude			
		Trying to be	We try to stay positive.	79-80
		positive		
			Even though I can see the changes	412-
			myself like but at the same time I'm	413
			hoping things aren't too bad.	
			But it's not easy seeing Chris the	94-95
			way he is but then I suppose it could	
			be worse. I don't know.	
			() and at the same time I always	78-79
			try that, em, we all try that	
			Usually we try to be in good form,	
			the kids and all that.	
		"Be positive	So let's say if I was someone who's	625-
		and make the	just starting to look after a person	638
		most of	with Motor Neuron Disease,	
		everyday"	would you have any advice for	
			me? What would you say to help	
			someone else to be prepared for	
			this role?	
			Just to be positive and to get up	
		"Carry on as	every day and make the most of	
		normal"	every day and to carry on as normal.	
			()	
			I would say carry on as normal once	
			you get over the initial shock and	
			make the most of every day and	
			don't let anything pass by that you	
			could do.	

Note: More on the PEG under Dysphagia theme

			A 1 1 2/1 1 2/1 1 1	C 4 0
1		Appreciation	And don't be, don't be sad over it.	640-
		of the "golden	Like be sad, okay you'll cry and cry	643
		time"	but generally make the most of it,	
			make the most of it. A cousin of	
			mine said to me I don't want to see	
			you crying. This is your golden	
			time. You enjoy it and she said	
			you've plenty time for crying after.	
			(Sounds upset).	
		Hope for slow	I just kind of hope that like when he	405-
		progression of	does the breathing there's so many	410
		MND	different things that they'll be kind	
			of I just hope that there won't be,	
			dis-improve too much. Doctor	
			usually like examines his lungs and	
			I just hope like that he doesn't, they	
			don't have anything too bad to say to	
			us.	
		Separated	I found nothing like but and I asked	639-
		MND from her	everybody, everyone I ever met	640
		home life	questions, loads of questions and	
			questioned everything.	
			Do you mean like?	
			The experts.	
			The experts, yeah.	
			And but then I put it behind me	
			when I'd come home again and I'd	
			just carry on as normal.	
	2.2. Seeing	Hard to be	$(\dots)$ we just tried to stay positive and	143-
	deterioration	positive	we still do but it's harder because	144
1				
			Chris is deteriorating.	
		Aware of	() even though it would be a huge	177-
		upcoming	() even though it would be a huge shock to change our lifestyle as	177- 179
			() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we	
		upcoming	() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows	
		upcoming	() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to	
		upcoming	() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows	
		upcoming deterioration	() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()	179
		upcoming deterioration Noticing	() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college () Like he went to Vienna last summer	179 210-
		upcoming deterioration	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs).</li> </ul>	179
		upcoming deterioration Noticing changes	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> </ul>	179 210- 211
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then</li> </ul>	179 210- 211
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while and I don't know whether that's it or</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I though the took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its because I'm at home and I see it all</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I though the took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its because I'm at home and I see it all the time in front of me rather than</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its because I'm at home and I see it all the time in front of me rather than when I'm at work I mightn't see it.</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its because I'm at home and I see it all the time in front of me rather than when I'm at work I mightn't see it. You don't know. I don't know but</li> </ul>	179 210- 211 203-
		upcoming deterioration Noticing changes Uncertain of	<ul> <li>() even though it would be a huge shock to change our lifestyle as well as Chris's but long-term we won't be able to do the cows because the kids will have to go to college ()</li> <li>Like he went to Vienna last summer and he walked Vienna like (laughs). He couldn't walk Vienna now.</li> <li>() before Christmas again I thought he took a step back and then he'd kind of go steady for a while and I don't know whether that's it or not. I don't know. Maybe its because I'm at home and I see it all the time in front of me rather than when I'm at work I mightn't see it.</li> </ul>	179 210- 211 203-

		sudden I don't know. He's not as	
		well as he was like.	
	Monitoring	I'm getting more worried about	319-
	deterioration	Chris all the time so I take out my	331
		phone and I take down the results of	
		the snip test that they do and, em, I	
		suppose I like to keep my own	
		record of how he's doing, in my own	
		mind. I don't know but maybe other	
		people wouldn't be like that now.	
		Wouldn't want to do that but I was	
		sorry I didn't keep a record of how	
		well he was last, you know, how he	
		does in these little tests that we do	
		when we go up. That's all I can	
		think of but I have started doing that	
		now myself that I keep the results of	
		the tests now, how he's improving,	
		dis-improving, em, oh sure I can't	
		really say.	
	From being a	He was always the leader on the	607-
	leader to being	holiday. That was a big change	609
	cared for	[having to be cared for]	
	Living day by	Are you planning to go somewhere	610-
	day	this year?	613
		We have no plans. I don't know. Its	
		more day-by-day this year	
		definitely.	
		Yeah.	
		Because there's bigger changes and	
		it will depend on this tube and lots of	
		things like.	
2.3 Emotions	Frustration	We always meet the same answer	332-
and feelings		that they can do nothing for us like	341
		basically, you know, I don't know	
		there doesn't seem to be any answer	
		You're hitting a brick wall the whole	
		time like but, em, it doesn't make	
1		time like but, elli, it doesn't make	
		any difference. It doesn't make any	
		any difference. It doesn't make any difference because nobody can	
		any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of	
		any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going	
		any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody	
	Lack of help	any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything.	430
	Lack of help for her	any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything. I don't mind going anywhere that	430
	for her	<ul><li>any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything.</li><li>I don't mind going anywhere that will help him like but I don't see the</li></ul>	430
	-	any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything. I don't mind going anywhere that	430
	for her husband	any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything. I don't mind going anywhere that will help him like but I don't see the point of course.	
	for her	<ul> <li>any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything.</li> <li>I don't mind going anywhere that will help him like but I don't see the point of course.</li> <li>I suppose I'm getting more worried</li> </ul>	430
	for her husband	any difference. It doesn't make any difference because nobody can suggest the slightest thing that's of help. That's why we gave up going to [name a service] because nobody can do anything. I don't mind going anywhere that will help him like but I don't see the point of course.	

F	Financial	I suppose financially I'm bringing in	179-
	oressure	the money at the moment because	181
P P	jiessure	there isn't any profit in [husband's	101
		work] at the moment.	
	Future as	() even though it would be a huge	177-
	'huge shock''	shock to change our lifestyle as	179
	nage sho en	well as Chris's but long-term we	1.72
		won't be able to do the cows	
		because the kids will have to go to	
		college ()	
	Constant	I am conscious of the background	171-
a	awareness of	all the time but I still enjoy very	172
	MND	much what I do [her work] but, em,	
		I suppose it a I don't know.	
	Self-blame	I suppose I could feed him better too.	475-
		() sometimes I'd be beating	478
		myself up for that [low intake] but	
		sure, you know, he has very good	
		food.	
	Coping	Well I suppose I find I can kind of	144-
		cope away with Chris and life but	146
		then when something else happens	
		anyone that's when it knock you	
		big time. If you understand? I	
		don't know ()	
0	Constant rush	And usually we're racing (laughs),	521-
		running around. Well Chris isn't.	523
		He spends of his time sitting here	
		now but we, em, we're all running	
		around him so like going, here,	
		there, doing everything but when	
		you go now the match will have to	
		be sorted	
	Being busy all	() I just watch Fair City and	75-76
t ti	he time	that's just purely for a break I	
		suppose because it's always very	
		busy ()	
	Being	() when I'm at home I live just	169-
	completely	totally separate lives when I'm at	170
	mmersed at	home. I'm completely immersed in	
h	nome	home. Then when I go to work.	

	Proudness	And all the children are brilliant.	659-
		They just, anything at all done.	661
		They're just And they were	
		never like I read somewhere that	
		God they'll be ashamed and	
		embarrassed about it. They're	
		never ashamed or never	
		embarrassed. They just go with it.	
		When you live with someone you	
		have a very clear, you can really see	
		like how difficult things are and	
		they're very good. They make	
		every effort to, em, to help him.	
	Desire to	I don't know should we be talking	644-
	protect her	more about it with them. I don't	646
	children	know because you can't hide it like	
		and I know that they, they know	
		everything about it like more or	
		less.	
		The children don't know about the	330-
		tube yet.	331
	Loving her	How do you enjoy your work?	19-21
	work	Yeah I love my work. I love my	
		work life. That's just it. I love my	
		work. (Laughs).	
9			

Jackie tries to maintain positive feelings and hope. She would like to protect her family from various consequences of MND as much as she can. She tries to bracket the disease from her family life by "carry on as normal". She tries to be positive (note that she uses the word "try" a lot here). She may be aware that this is not fully achievable and all she can do is "try". She knows that her husband will not live for a long time and she feels she should appreciate every moment spent together. She knows that the grief should be left for after her husband is gone. Now is the time to enjoy life (try to enjoy). It seems like a struggle between emotions and pragmatic reasoning. It may sound logical to use the time to enjoy life and not to get upset while Chris is still alive and reasonably well. However, she knows that Chris's death in unavoidable and that she will be left alone with children and a [anonymised], which she would have to sell at some stage. It is worrying – seeing her so emotionally drained and exhausted now while she is trying to be positive and strong. How will she be when she is alone? How will she be when she lets her grief out? One can only hope that her love for their children will keep her going and give her strength.

MND is the only disease known to me that "by right" you should have no hope. There are treatments for many other serious diseases (cancer). They may/ may not work, but they are available. Other progressive neurological diseases can have periods of remission (MS) or can progress over a long time (Parkinson's disease). MND is cruel, rapid and has no cure. This is typically made clear to any patient at the time of the diagnosis. However, Jackie still has hope. Her main hope is for the slow progression of MND. She knows her husband is going to get weaker but she wants to delay this as much as possible. She is accepting the way he is now and wants him to stay like that forever. She seems happy with what he can still do and the fact that his mobility is still functional and he is independent. When they go for medical appointments Jackie hopes to hear good news, which for her would be little or no deterioration from the previous visit.

This is a very challenging role to play and emotionally draining. Jackie carries a lot of hidden emotions and it can be exhausting. She burst into tears the minute she sat down to talk to me and it

didn't stop. She kept saying: "I am fine normally. It is only when I talk about it. I don't usually talk about it. I am fine".

While trying to be positive and carry on as normal Jackie can see the deterioration of her husband. He is physically weaker and unable to participate in family life as he used to. She cannot deny it as it is happening in front of her eyes. MND is visible to everyone. This is very hard: to produce a smile on her face while her heart is breaking with pain. There are changes in their life caused by the progression of MND and Jackie may be attempting to hide them or to make them less significant/ impactful by trying to be in good form. This must come at a high price.

At the same time Jackie is full of frustration. "You're hitting a brick wall the whole time like but it doesn't make any difference". All the medical appointments achieve nothing for her. No one can help her husband to recover. She is trying to keep him well but she know it makes no difference in the end. MND always wins. She looked for answers herself (by checking the web sites) and she asked the experts. She got no answers.

In relation to the swallowing: Jackie tries to do her best to ensure the food is high in calories and safe in consistency. She spends time thinking about what to buy and how to cook it. At the same time she knows no consistency is fully safe, so her efforts are failing (He, em, like when he eats, even like if you have everything mashed up he, em, like he'd caught, it gets caught and it's not easy). She is not giving up, but she is blaming herself for not being able to do better (I suppose I could feed him better too. I don't know).

All above factors place Jackie on edge. She can barely cope with her everyday life. If something additional happens it "knock her big time". (Well I suppose I find I can kind of cope away with Chris and life but then when something else happens anyone that's when that it knock you big time. If you understand? I don't know (...).

Jackie is also physically exhausted. She is not sleeping well, she works full time, she has additional [anonymised] duties, she has four children and a sick husband. She is neglecting her own needs to look after other members of her family. She gets no break. She also is the only earner in the family now, since Chris is unable to do the work. Although she is ready to give up work for Chris she knows that this would put financial strain on the family. Jackie is worried about her future. The

[anonymised] would have to be sold at some stage although she is going to delay it. She is going to be left with the kids. The family is so important for her and for Chris. Now MND is going to damage that forever. The kids are going to start college and soon she will be living alone. This would be a massive change for her, something she would never have wanted to have.

Jackie's priority is her family. This is her source of happiness and pride. She admires the children for the way they accept their father and how they are trying to help her every day. She would like to protect them from additional stress and she is not sure how much information to give them. MND cannot be hidden so the kids can see the problem, but still they may not be as aware. It is hard to know how much information to disclose and when. Jackie is a part of her family and almost doesn't feel her own identity (She says "we" a lot). Jackie is completely immersed in her home life. She loves her job and she acknowledge that it is a break for her too (although the MND is all the time in the background). However, she is ready to give work up in the morning at Chris's request. I see this as another example of neglecting herself – not only does she need work for her mental wellbeing and sense of satisfaction, but she is the only person bringing money to the house. Giving up work may mean getting loans which she would have to pay back in the future (unless they sell the [anonymised] which may be a solution, but even at that she would sacrifice herself. There will be no distractions from MND anymore). Jackie's priorities are clear. At the moment she has two separate lives and that helps her to cope. She is ready to change that to be with her husband as much as possible and as much as he requires.

I think it is very important for Jackie to have her two separate lives. Even though she is aware of MND all the time, work brings her many positive aspects (joy, satisfaction, fulfilment). "It is a break" she says. A break from frustration, injustice, pain, grief, and no hope. It seems important to have that second life, a moment of disconnection, a time when she is surrounded by her old colleagues and where MND is not the most important part of the day.

3. Dysphagia	3.1. The visibility of dysphagia	Different diet	() and it would all have to be, em, like it would, yeah, obviously it would be different to everybody else's on the table.	244- 246
		Signs of dysphagia	How did you notice that the swallow was off a little bit? Oh he wasn't able to eat it like. You notice like if somebody couldn't eat it like.	278- 283
		Children able to see signs of dysphagia	Okay, okay and were you talking to the kids about the swallow at all? Well they can see that. They can see that.	125- 135
		Visible signs of dysphagia	They know, they can see that because Chris eats slowly number one, and number two he, em, can kind of gag on his food at times, and he can get dreadful coughs trying to bring up stuff and they can see that his dinner is soft and that his potatoes are mashed up with butter and they can see.	
		Coughing during meals	And what does he do if he's eating and if he starts coughing? It depends. It depends sometimes he'll go outside the door for air and if he's here we'll open the window for air or sometimes he'll just take it out of his mouth and put at the side of his plate if it comes up. Does he continue eating then? He continues eating.	313- 314

It is interesting to note how Jackie and her children see swallowing problems. They are focused on how Chris eats so they can see that his mealtimes are longer, that his food looks different and sometimes he coughs up/gags on food. He may need to open the window to get more air when eating. They are focused on him so they can see that. This was not as evident in any family so far.

3.2 Managing	New approach	I approach the food differently.	222
dysphagia at	to food		
home			
	Concerned	Definitely yeah. When I go	218-
	about nutrition	shopping I think of calories most of	220
		the time	
		Well I buy cream and double cream	213-
		on a very regular basis.	217
		That's for calories, is it?	
		Yeah for calories yeah.	

Having	And would you keep it different	943-
unhealthy	now for the kids or do they all	947
food at home	have the same?	2.1
	No, he has his own food. They're	
	very health conscious the four of	
	them.	
	Are they?	
	They are very, they are. They all	
	play sport and they wouldn't dream	
	of eating any rubbish.	
Eliminating	Is there any other food like that	270-
unsafe foods	that he's not that is off the list at	272
	the moment?	
	Well anything hard (laughs).	
	Anything hard, okay. So it could	
	be meat, it could be fruit, yeah.	
	Oh fruit, apples and all that is gone.	
	Okay.	
	Yeah, yeah, unless stewed apples	
	but	
Modifying	Yeah dinner then would come then.	239-
food textures	I would basically, em, we have	244
	mince now today. Well basically	
	then I would mash the, I did mash	
	Chris's potato with butter and make	
	it butter and creamy and the	
	vegetables would be mashed up with	
	plenty butter (laughs) added as well	
	and the meat would be obviously	
	soft. Em, it would all be very soft	
	()	
	White sauce, we'd use white sauce	929-
	and gravy and he has ketchup. He	932
	used never eat tomato ketchup much	752
	or we'd have YR sauce but he	
	always has sauce yeah and I'd buy	
	new products like, you know, bacon	
	and ham, you know, a different	
	sauce that I could put over	
	something yeah	
Supplements	Well I suppose I know he's having	1158
Supplements	the Fortisips so that gives me	1150
	assurance.	
Thickened	Thickener alters the taste	274-
liquids	What way? Do you just mix it	274-277
(Preparation)	with a spoon or what way do you	<i>211</i>
	do it?	
	I put it in and mix it with a spoon for	
	twenty seconds as it says on the tin.	
	Well it went lumpy at the bottom so	
	I obviously didn't mix it enough that	
	day and he tasted but he didn't get	

	far in the cup. Like he didn't get as	
	far as the lump in the bottom to taste	
	it. (Laughs).	
	(Laughs). He knew sooner.	
	He knew straightaway. He said the	
	milk is gone off and I said no it's not	
	but anyway	
Husband's	He doesn't like the thickener, he just	269-
dislike	doesn't like it. He can taste it.	273
	Has he tried it for a bit?	
	He's tried it yeah. Maybe I put in	
	too much. (Laughs)	
Adding	I must kind of cut it down again and	281-
thickener	put in a little bit and maybe he	284
covertly	mightn't notice it.	
	So do you put it in yeah?	
	Sometimes I put it in but he has told	
	me when I put it in. (Laughs).	

Jackie's attitude to food has changed and she is admitting it. Her food choice is now dictated by high calories and soft consistency. This is all new for her as they used to eat very healthy. Now she has "rubbish" food at home and it is unusual. All healthy eating priorities are not important as MND has its own rules. I remember people in the clinic worrying about raising their cholesterol. MND wins with that. You have some changes i.e. longer survival when you eat a lot and keep your weight up (this again is a fight you are not able to win anyway). Jackie seems very proficient at modifying food consistencies. She does it by longer cooking, chopping, mashing, adding extra sauces. Jackie is very aware of various ways of modifying food consistencies to make them easier to swallow. She provided information clearly and it was almost effortless. The diet modification techniques are an integral part of her life now. I wonder if her attitude towards food has changed for her. Chris dislikes thickener but Jackie feels she must keep adding it in. She blames herself for his reluctance to take thickened fluids ("Maybe I put in too much"). She thinks about adding the thickener behind Chris's back. She must know it is there to protect him from aspiration. Although she respects Chris's decision and choices her main priority is to keep him alive, so would do anything to achieve that.

uchieve inui.	achieve mai.					
	3.3 Chris'	Decreased	It was mince and vegetables and em,	270-		
	changed	appetite	for tea now tonight he might have	274		
	approach to		lasagne or he might have, em,			
	food		actually there are more and more			
			things he doesn't like. He used to			
			like fish pie and he doesn't like that			
			anymore (laughs).			
			So that's changed as well yeah.			
			He doesn't like as many things like.			
			Is it harder now to find something	275-		
			to cook?	277		
			It's harder to find things yeah and			
			before he goes to bed then he'll			
			have crackers or biscuits again so			
			biscuits and tea or coffee, a latte.			
			How's his appetite?	255-		
			It can be good some days but	259		
			generally he doesn't have anything			

			like the appetite he had before. I'd	
			say he eats for the sake of eating,	
			you know. He feels he has to eat.	
		Becoming a	Yeah, I would always think of	909-
		"fussy" eater	what's soft and like I'd put down,	912
			em, often put, em, if it was lamb	
			chops or I was looking for	
			something nice, you know, he's not,	
			he was never a fussy eater but he's a	
			bit more fussy now so like I'd put the	
			chops into the oven at a low heat for	
			a very long time and I would totally	
			change everything.	
		Variable	I suppose he's better some days than	948
		intake	other days.	
	3.4	Dysphagia is	How aware do you think people	454-
	Understanding	like "every	are, generally in society, of	463
.	of dysphagia	issue in life"	swallowing problems?	
			Unaware (laughs).	
			Well I suppose like I was unaware of	
			[confidential] or anything until it	
			happens to yourself.	
			That's the way with every issue in	
			life. Like parents in work they're	
			unaware of things until it comes to	
			their own door.	
			Everybody has something.	
			And you have to go down So it's	
			swallowing. Yeah that's the way it	
			is.	
		Awareness of	You cook the same meal yeah you	251-
		the	just add more butter and cream to	253
		progressive	Chris's.	
		nature of	Yes and have it mashed. I wouldn't	
		dysphagia	say pureed because it's just mashed	
			at the moment with a fork	

Dysphagia in MND is progressive. Every disability in MND is progressive. You may attempt to solve a problem but every solution is temporary. Jackie adapts to Chris's changing needs. His appetite has dis-improved and he does not take pleasure from eating anymore. She tries to solve that by researching new foods, new cooking ideas. She gets a sense of security from having nutritional supplements at home. This is one of her aims to keep her husband alive (ensuring adequate intake). She knows that dysphagia will progress and she feels prepared to manage that. She knows how to adapt food consistencies and use supplements. She will not be looking for professional help as she believes the services don't really offer any solutions.

Jackie respects Chris's choice regarding food. He stopped eating bread after a bad choking experience. However, when they went away he attempted to eat bread in a restaurant and she didn't stop him. instead, she accommodated his wish by:

"I just cooked one side of it and em, I put a lot of butter on it and put a bit of marmalade on it and he'd try it and he didn't anything, like the crust and all were cut off but he had a bit of it that day". I think Jackie is doing outstanding work in relation to the management of dysphagia. She is realistic and pragmatic, yet accommodating and gentle. She has no previous experience of dysphagia at all: **Did you have any knowledge of swallowing before Chris** ...?

No.

**No.** (412-414)

Jackie seems not to be particularly focused on dysphagia. This is just one of the steps to help her achieve her goal: keeping Chris alive. She says that dysphagia is like "every issue in life". You will be unaware of it "until it comes to your own door". She accepts it as an unavoidable feature of MND and she feels able to manage it. "Everybody has something" she says. **Yeah.** 

And you have to go down... So it's swallowing. Yeah that's the way it is."

There is not much emotion associated with talking about dysphagia in particular. Dysphagia is another problem she needs to manage, but she is looking beyond the swallowing in order to achieve her main goal.

goai.			1	
	3.5 Non-oral	"Monumental	And here we are now six weeks	449-
	feeding	decision"	later and after coming to this	450
			monumental decision in our lives	
			we're still waiting.	
			He took his time with us [a doctor]	97-98
			and I suppose, it's up to yourself to	
			make the decision but it's a very	
			tough decision.	
		Consulting	I suppose that should be the next	118-
		others	step now. When you talk you kind	119
			of get ideas yeah because Henry	
			[other man with MND and PEG]	
			did email him two or three nights	
			ago but he didn't mention the tube.	
		Conflicting	The nurse was here then and, em,	335-
		opinions from	she said you should go up to see our	344
		professionals	doctor. She's always dealing with	011
		proressionais	the tubes. So we went up there.	
			That didn't Oh I suppose you	
			make up your own mind like	
			basically on things. We were down	
			at his physiotherapist and em, we	
			were telling her and what did she	
			think of the tube? We were asking	
			her and she said yeah. Oh look	
			there's one nurse in here, now she	
			constantly deals with the tube. I'll	
			bring her out. She came in and she	
			doesn't want to commit either way	
			but they weren't coming out very	
			strongly in favour of it as I would	
			like, you know, so kind of Chris	
			said no.	
			And how much do you know	422-
			-	422- 428
			about looking after the tube then	420
			and the whole feeding thing?	

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		I don't know anything much about	
		it. They say it's Well it depends	
		who you speak to. Some say it's very	
		straightforward, others say it's not at	
		all straightforward. One person says	
		it's as easy to get a chest infection	
		from a tube as it is to get from	
		swallowing.	
		Okay.	
		So I don't know. I don't know	
		really	
	PEG	And how did he decide to go for it?	327-
	recommended	How did he decide? Em	330
	by	Like has he seen the tube before or	
	professionals	how?	
		No, they all recommended it. They	
		recommended it.	
		The [anonymised] seemed to be very	78-81
		much in favour of it like. They seem	
		to be in favour of it.	
	Feeling	So do you feel you're getting kind	82-86
	confused	of contrasting messages or is it	
		that it's just so complex?	
		I don't know. I really don't know	
		because it's been brought to up, it	
		has been brought up to us a lot in	
		the past for the peg and now when	
		we're kind of going about it like we	
		don't know what we're doing.	
	It's Chris's	A lot of people were saying it to	26-30
	decision	me, do you know that it probably	
		would be better if he got it () but	
		Chris wasn't in favour of it so we	
		left it.	
		So it's basically back to Chris	76
		whatever he decides.	
		Well its Chris's decision.	86-87
		I left it to Chris. I can't decide	
		whether he gets a peg or not. It's	
		whatever Chris wants	
		It's up to Chris. I hope he'll get the	103-
		strength to make his own decision	104
		because I wouldn't push it.	
	Husband not	How it happened, em, well, em,	20-24
	anxious to get	about this time last year the subject	
	the PEG	was first broached with us that	
		maybe he should go for a peg and I	
		think it might have been said in	
	Respecting	[hospital] first and then we went to	
	husband's	[another hospital] and it was said in	
	choices	[there] as well and then, em, Chris	

	wasn't anxious for it and I would go	
** 1 1 1	with whatever he wanted.	222
Husband slow	They recommended the tube and	333-
to decide	em, he was very slow to accept it or	335
about the PEG	to take it on board that he might	
	need it or want it so he said no for a	
	long time and em, we looked up a	
	lot about the tube.	
Attempted to	One day coming down in the car	344-
talk to the	then I said like the speech therapist	347
husband about	took out the tube again to Chris and	
the PEG	I said what did she say about the	
	tube Chris and he said leave it all	
	back there (laughs) and I said no	
	more.	
Husband had	Chris's father died last year and I	348-
negative	know it's totally different now. He	354
experiences	had, em, oh he was continually	
with other	getting urinary tract infections so he	
types of tubes	had a tube inserted down there and	
cypes of tubes	oh infection, infection, infection	
	from the time he got the tube to the	
	day he died. So he was allergic to	
	tubes. He didn't want to hear tubes	
	at all. I suppose Chris was his	
	primary carer even though Chris was	
	very sick, he was the only person	
	that his father, you know, he wanted	
	Chris around all the time.	
Would like	So then eight weeks ago we, Chris's	30-35
him to get the	aunt came down and she spoke quite	
PEG to	forcibly about it and I said and then	
prolong his	a dietician had said it to us as well	
life	and the speech therapist said it to us	
	in [hospital] like, well Chris what do	
	you think about the peg like? It	
	would prolong your life and it's a	
	good thing and that so then anyway	
	about eight weeks ago we decided	
	that, well Chris decided, he said that	
	I think I will email [hospital] about	
	the peg, so we did.	
	Oh look there's one nurse in here,	343-
	now she constantly deals with the	348
	tube. I'll bring her out.	
	She came in and we knew her	
	because she used to be a neighbour	
	that lived here before. I didn't	
	really know her but Chris knew her	
	and em, a very nice person and	
	obviously she doesn't want to	
	sectoring one doesn't want to	

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		commit either way but they weren't	
		coming out very strongly in favour	
		of it as I would like, you know, so	
		kind of Chris said no.	
		A very hard decision. She gave, I	71-73
		thought going up that day that, em,	
		oh great now she'll tell him get the	
		peg and all will be great and but	
		anyway no she wasn't She didn't	
		push the peg at all.	
	Torture of	What's happening with the peg	322-
	waiting for	tube?	326
	gastrostomy	I don't know. Its five weeks now	
	gubulostomy	since we broached that subject. I	
		rang Holy Thursday.	
		So you're waiting to have it done?	
		Yeah, yeah. Its five-and-a-half	
		weeks now. Today is Tuesday	
		again. Now is it six weeks today?	
		Yeah its six weeks.	
		() We had emailed them two	269.0
		weeks prior to that that, em, please	368-9
		put everything in motion.	4.40
	Aware that	And I'm also very aware of the fact	440-
	husband may	that after a certain stage you can't	442
	become too	get the tube like. So I actually, I	
	unwell to get	don't know. I worry sometimes is he	
	the PEG	still alright to get it but I hope he is.	
	Anxious to get	But like I did ring and suppose I be	444-
	the PEG	ringing again today. I didn't ring	450
		now yesterday but it is on my mind.	
		(Long pause). I thought really when	
		he made the decision like because	
		there was so much, people saying to	
		us like you should get the tube, you	
		should get the tube. I thought that	
		when he decided that he would get	
		the tube. And here we are now six	
		weeks later and after coming to this	
		monumental decision in our lives	
		we're still waiting.	
	Told that PEG	She says it brings as much	73-72
	can be	problems as solutions. You can get	
	problematic	infections, chest infections from	
	· · ·	having the peg as well as from	
		having it and the food can go up as	
		well as go down and she wasn't	
		pushing it at all as was the man	
		yesterday.	
		Yesterday he said that it will not	52-62
			52-02
		prolong life.	1

1	The transfer of	It might agag your situation but it	
	The trauma of	It might ease your situation but it	
	unexpected	will not prolong life.	
	appointment	He spoke of the ethical side of it as well.	
		And what did he say about that?	
		Well he just said he gave us his	
		own background. That he's written	
		a I haven't read them but he's	
		written a lot of papers and things	
		from the ethical side like, like of	
		people, I suppose. He would be	
		quite willing to give Chris the peg	
		like with older people and things	
		like that, you know, but, em, we	
		would have been aware of all that	
		side of things. We would have	
		discussed that with people. We	
		would be aware of that and we had	
		met another nurse and she spoke	
		about ethical issues as well	
		(laughs).	
		It's whatever Chris wants but like	87-88
		that was only yesterday. It's only	
		sinking in. I haven't spoken to him	
		today.	
	Having to re-	Right and how did it go	732-
	consider	yesterday?	734
		Em, well there were no answers, you	
		Em, well there were no answers, you know, it's up to Chris to decide	
		know, it's up to Chris to decide	
	"crossing a	know, it's up to Chris to decide	418-
	"crossing a line"	know, it's up to Chris to decide again now.	418- 419
	-	know, it's up to Chris to decide again now. He feels it's crossing a line and he's	
	-	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot	
	line"	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything ()	419
	line" PEG as	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I</li> </ul>	419
	line" PEG as additional	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make</li> </ul>	419
	line" PEG as additional	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough</li> </ul>	419
	line" PEG as additional	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> </ul>	419
	line" PEG as additional	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> <li>Very tough in many ways because</li> </ul>	419
	line" PEG as additional duty	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> </ul>	419
	line" PEG as additional duty PEG as	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> <li>Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush</li> </ul>	419
	line" PEG as additional duty PEG as support	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> <li>Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every</li> </ul>	419
	line" PEG as additional duty PEG as	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the	419
	line" PEG as additional duty PEG as support	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when	419
	line" PEG as additional duty PEG as support	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> <li>Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's</li> </ul>	419
	line" PEG as additional duty PEG as support (dilemma)	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's just tough. I don't know. I didn't	419 97-104
	line" PEG as additional duty PEG as support (dilemma)	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's just tough. I don't know. I didn't I'm hoping this operation, now this	419
	line" PEG as additional duty PEG as support (dilemma) Fear of the future with	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's just tough. I don't know. I didn't	419 97-104
	line" PEG as additional duty PEG as support (dilemma) Fear of the future with PEG	<ul> <li>know, it's up to Chris to decide again now.</li> <li>He feels it's crossing a line and he's well educated and he knows a lot about everything ()</li> <li>He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision.</li> <li>Hmm</li> <li>Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's just tough. I don't know. I didn't</li> <li>I'm hoping this operation, now this big operation won't set him back.</li> </ul>	419 97-104 415
	line" PEG as additional duty PEG as support (dilemma) Fear of the future with	know, it's up to Chris to decide again now. He feels it's crossing a line and he's well educated and he knows a lot about everything () He took his time with us and I suppose, it's up to yourself to make the decision but it's a very tough decision. <b>Hmm</b> Very tough in many ways because like you, okay Chris's walking around and he doesn't have to flush out anything or do anything every day to himself like he's and at the same time then if you get sick when you need something to feed you, it's just tough. I don't know. I didn't I'm hoping this operation, now this	419 97-104

don't know what's coming down the
track like.
Hmm Would you change
anything?
I don't know. I don't
Like I don't know. I just don't know
what to expect. He can just have it
there and not use it but also like he
would be so tired, you know, he was
told he'll have to be sitting. He can't
use at night unless he's
Sitting upright or?
He was saying he might use it during
the day preferably, em,

It was exhausting to listen to Jackie talking about the gastrostomy tube. For her, it was the hardest decision ever. It was a process of consulting others and getting conflicting advice, being aware that it is a way of prolonging Chris's life (so achieving her main goal) while at the same time wanting Chris to decide for himself (respecting his decisions). Consulting others made the process particularly complex and confusing for Jackie. She was hoping for everyone to encourage Chris to get the gastrostomy tube, but not everyone did. She stayed silent for as long as she could, but when she noticed further deterioration of Chris's swallow she gently started to talk to him about non-oral feeding. She couldn't wait and see him wasting away and missing the opportunity of having the gastrostomy tube (while she believed it will help him to live for longer time). The decision was huge. Monumental. Chris had some bad experiences with other types of tubes, which caused him to dislike tubes in general. However, one day he said yes and Jackie acted immediately. She rang and emailed the hospital a number of times and... nothing happened. For over 6 weeks they were waiting for an appointment. The time was long enough for Chris to change his mind or to deteriorate beyond the point where he would be still a candidate for this procedure. Jackie was stressed out. Unexpectedly, a doctor who was supposed to call them for the gastrostomy procedure called them for a consultation instead. He was not in favour of non-oral for Chris. He gave his reasons. He quoted his own research. He smashed Jackie's hope ("It will not prolong life he said"). He asked Chris to reconsider it for two more weeks. Like he hadn't considered it for over a year. Moreover, the team of MND clinicians seemed to be in favour of non-oral feeding too. Jackie was traumatised and exhausted after that appointment. There was less animation on her face, less emotions in general. She was completely overwhelmed and concerned. She was delighted when Chris consented for gastrostomy and now the process of deciding was to start again.

I felt absolutely shocked after hearing that story. These were not naive people hoping to win with MND or people who were not aware of the consequences of living with non-oral feeding. They were aware of the ethical side and every possible medical complication. That doctor hadn't met them before. He knew the disease, but not how the disease affected Jackie and Chris and how they coped with it, adapted to it, lived with it. Looking at Chris's type of MND and his stage, the choice is to either get the gastrostomy now or to not get it and stop being able for any oral intake soon. This could kill Jackie. This will become a problem for all when the medication can't be swallowed or when Chris is dehydrating. There would be nothing that Jackie could do to ease the situation as the oral route will not be an option anymore. She needs the security of the PEG, although she is aware that it is "crossing the line" and it is like a symbol of losing his independence / losing their "normal life" forever.

3.6	MND Clinic	And how happy are you to go in	307-
Professional		the room and have a few	309
Services		professionals sitting there?	
		It's fine. () its better.	

		Yeah, I think it's nice. () I would	
		have no issue with that. It's	
		probably better actually. I think it's	
		actually better that they're all	
		They all know everything about you	
		together.	
		I don't like, em, I think it's a good	
		system. I don't like to be critical.	
		I don't like to be critical so I don't	
		like saying anything bad but I know	
		in () now it's a bit more one-by-	
		one and I think it's better together	
		because they're all informed about	
		everything.	
	Training	I just go to the workshop, like the	332-
	"Came home	MND swallowing workshop around	345
	upset"	that. I was on my own.	
		And what did you think of it?	
		It was fine. I suppose I had already	
		been to the swallowing, you know,	
		I'd spent three quarters of an hour	
		above with Chris so I don't think I	
		learnt anything new. ()	
		Because I had listened to it all and	
		been through it all.	
		()	
		They went through the four types of	
		food like, you know.	
		All the diets.	
		The diets and they went through the	
		thickening, the thickener and they	
		went through	
		They had cups, you know, and they	
		had different cups you can hold. I	
		suppose I wouldn't have been aware	
		of those now to be honest (laughs)	
		and we hadn't got to that stage yet.	
		Hmm	
		And, em, (short pause) well to be	
		honest most of it I felt I hadn't learnt	
		much new there because I had been,	
		I spend an hour; I had been with	
		Chris the previous week or two. I	
		had been above there on an	
		individual basis and I kind of	
	Didn't identify		316
	Didn't identify	There were only two others ()	346-
	with other	I just felt I hadn't much in common	367
	caregivers at	with them.	
	the workshop	Right.	
		I hadn't much in common.	

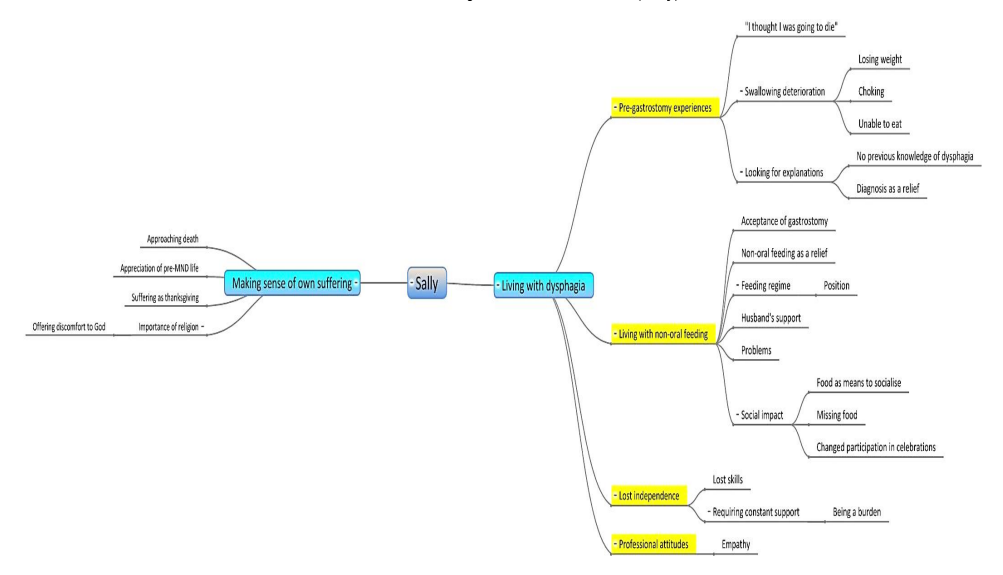
Waste of time	So, I don't know. I was there that	382-
	day now without Chris and I didn't	405
	feel any better.	
	I felt it was a waste of my time being	
	away from Chris to be honest. ()	
	Yeah, next question (laughs).	

Jackie's relationship with the health services is interesting. She attends them with Chris but she doesn't believe that they can offer any help. She attends them to monitor Chris's deterioration as she is unsure of her own observations. She prefers to be seen by the team together, this experience does not seem to intimidate her. She sees it as a better usage of her time.

The swallow workshop she attended was not a positive experience. First of all it was a repetition of the knowledge she was already given at the SLT consult. That was the way she perceived it anyway. She noticed various drinking utensils and she knows that Chris may need one in the future. This was new knowledge for her and maybe there were other examples which she didn't report to me. What really complicated the experience for Jackie was hearing the experiences of other caregivers, some very traumatic for her. She came home upset and she felt she "wasted her time being away from Chris". Was that something the SLT should/could anticipate? It is still possible that Jackie feels so confident in managing dysphagia due to attending consultations and the workshop and getting a good understanding of dysphagia from the professionals involved? It is simply not possible to ignore the emotional side of her life. She was unable to appreciate hearing about various cups while at the same time she also heard that someone similar to Chris was almost dead.

It looks like work is the only place where Jackie immerses herself for 90%. When she is shopping, eating out, going on holidays and attending workshops the MND is in front of her eyes all the time. It changes the way she perceives and experiences the world around her. Only work allows her to escape.

#### **APPENDIX 8A: Mind Map – A Person with MND (Sally)**



#### Appendix 8b): Mind Map – A Caregiver (Jackie)

