IF YOU NEED FURTHER

INFORMATION

If you need more information or support, or would like to order our publications, contact our helpline.

PSPA Helpline 0300 0110 122

helpline@pspassociation.org.uk

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PSP

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LIVING WITH PROGRESSIVE SUPRANUCLEAR PALSY

PSP: what to expect from your care



PSPA is the only charity in the UK supporting Progressive Supranuclear Palsy (PSP) and Corticobasal Degeneration (CBD), which is sometimes known as Corticobasal Syndrome (CBS).

For more information about us, visit www.pspassociation.org.uk

You can use this guide to open discussions about your care with health and social care professionals and providers. This may help you access suitable support at the right time. The pages that follow highlight recommendations for good care, but you are unlikely to need all of these at once.

Use the questions provided to help you discuss your needs as they arise and plan ahead for future care.

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FOLLOWING

DIAGNOSIS

- Offered a follow-up appointment with a neurologist?
- Offered written information including access to supporting organisations?
- Offered a care plan and been involved in its development?

ORGANISING

YOUR CARE

HAVE YOU BEEN:

- Given the details of a named key worker/point of contact?
- Offered regular reviews of your care and medication, through your multidisciplinary team?
- Offered a referral for prompt management of symptoms by a specialist palliative care team?

MOVEMENT

AND BALANCE

- Referred to a physiotherapist and/ or an occupational therapist to help maximise your independence?
- Assessed and treated for risk factors related to your current and future needs e.g. osteoporosis, falls, visual disturbance?
- Assessed for equipment, adaptations and/or safety measures to meet your current and future needs?

VISION AND OTHER

SENSORY CHANGES

HAVE YOU BEEN:

- Referred to an ophthalmologist for assessment and possible prescription of specialist glasses?
- Referred for an assessment of your home environment to reduce any hazards?
- Given an opportunity to ensure any changes in vision are not affecting your communication?

COMMUNICATION

AND SPEECH

- Assessed by a speech and language therapist for your communication needs, or reviewed if your needs have changed?
- Referred for further assessment if you need communication aids?
- Given help to ensure any communication aids work well with other assistive equipment that you may use?

SWALLOWING

AND NUTRITION

HAVE YOU BEEN:

- Assessed and offered support/ treatment if you have saliva problems?
- Assessed for any swallowing, eating and drinking difficulties or associated issues such as chest infections or breathlessness?
- Given the opportunity to discuss alternative ways to receive nutrition, such as tube feeding?
- Given the opportunity to discuss and plan ahead for your future needs and wishes?

BLADDER AND

BOWEL

- Given advice from a dietitian on diet and fluid intake?
- Given advice on bowel care by the community nursing team?
- Assessed by the continence nurse and given access to appropriate products?
- Given a medication review if continence is a problem?

THINKING AND

BEHAVIOUR

HAVE YOU BEEN:

- Asked about any changes to your thinking or behaviour?
- Referred for a formal assessment if needed?
- Informed about access to emotional support for you and your carers?
- Given the opportunity to express your wishes in your care plan?

FATIGUE

AND SLEEP

- Referred to an occupational therapist for advice on fatigue management?
- Assessed and given advice if sleeping patterns are a problem?
- Given a medication review if sleeping is difficult?

PAIN

MANAGEMENT

HAVE YOU BEEN:

- Assessed by a physiotherapist if limb stiffness is a problem?
- Advised how to communicate that you have pain if your speech is affected?
- Referred to the specialist palliative care team to help with complex pain symptoms?

PLANNING

AHEAD

- Offered an early opportunity to discuss future care concerns about your condition and end of life, if wished?
- Given support and advice about planning ahead for future care?
- Given prompt access to specialist palliative care services, if needed?

WHAT CAN I DO IF MY

NEEDS ARE NOT MET?

If you do not receive the care or treatment you need, or have any problems, discuss this first with the relevant health or social care professional, or care provider.

If this does not improve the situation, make a formal complaint or appeal to the service involved. They should be able to provide information or advise how to do this.

IF YOU NEED FURTHER

HELP OR SUPPORT

PSPA resources for people affected by PSP: https://tinyurl.com/yxtevr4s

PSPA resources on benefits: https://tinyurl.com/yxaszfbj

PSPA resources on Continuing Healthcare Assessments (CHC): https://tinyurl.com/yxmq8yhr

PSPA resources on other sources of funding: https://tinyurl.com/y4wapz39

PSPA resources for Health and Social Care Professionals: http://tiny.cc/y1td6y

PSPA Research opportunities: http://tiny.cc/rmrd6y