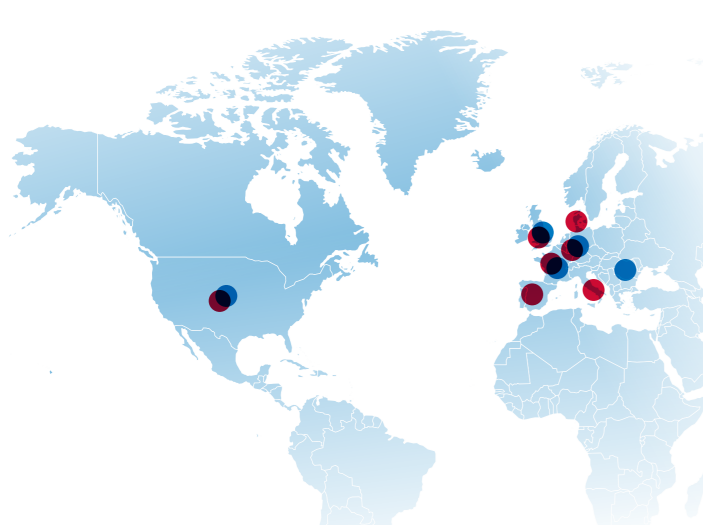






MS is associated with a high cost of illness and a clear deterioration in health and wellbeing of both people with MS and their caregivers.

There is a need to achieve better outcomes for people living with MS and evidence within the IMPrESS by the London School of Economics suggests that this can be addressed.



What is IMPrESS? The IMPrESS report assesses the

socio-economic and personal impact of MS and presents evidence on the need for a paradigm shift in the management of MS.1

Primary and secondary data sources used:

Primary data sources included: Adults with MS and caregivers

- (in France, Germany, Romania, UK and the USA) Clinicians (in Denmark, France, Germany, Italy, Spain, UK and USA)
- Secondary data sources included: Analysis of Health Technology Assessment (HTA)

recommendations

The IMPrESS report is an independent study from the London School of Economics, funded via a grant from Roche, that assesses the socio-economic and personal impact of MS and presents evidence on the need for a paradigm shift in the management of MS.



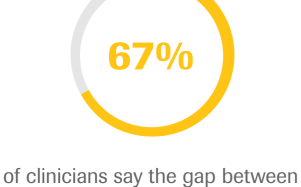
Improve quality of care and health outcomes. Diagnosis, treatment and management goals should be set

Early diagnosis and treatment are needed to secure

the best outcomes and should be routinely available

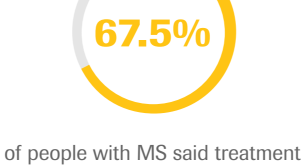
Delays in diagnosis can be minimised by making specialist care,

diagnostic and monitoring procedures widely available¹



than 2 months. People with MS said the delay was on average 1.9 years¹

diagnosis and treatment was less



should be started at clinical diagnosis; 31.3% however thought it should be first symptom¹



for public education around the symptoms of MS to minimise delays in diagnosis²

more effective disease-modifying therapies (DMTs) earlier

Clinicians should use **new** and

only considered in the presence of brain lesions, optic neuritis or severe initial relapse.1 57% of clinicians reported that they may delay switching to another DMT but it depends on the individual¹

92% of clinicians wouldn't start DMTs in a person with a

normal MRI, a tool used to measure disease activity. DMTs

monitoring regimens and the use of MRI in capturing disease activity

Further research is needed on the effectiveness of more intense

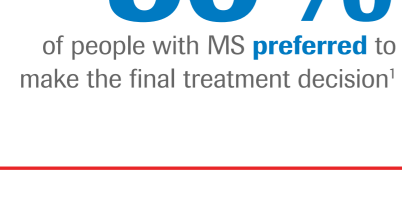






Engaging the individual with

Actively involve people with MS in decisions about disease management





increases satisfaction¹

MS in decisions provides

health benefits and



Robust evidence needs to be generated HTA recommendations differ

Implement informed decision making

National registries should be in place and their data routinely used

In the UK there is no complete

registry of people with MS,

across countries

of the MS population¹

but in Germany databases exist which cover

There is a need to standardise the approach when including the views of

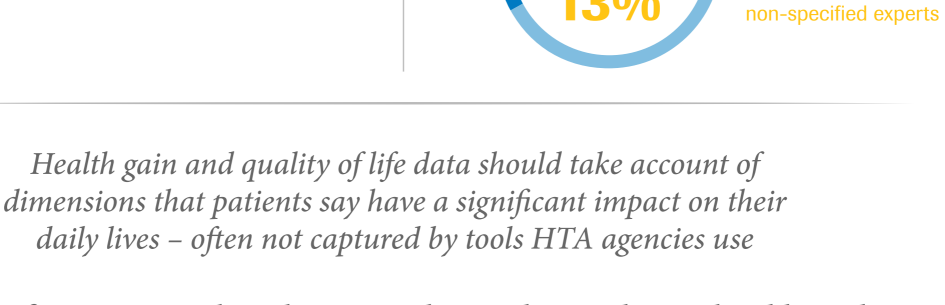
people with MS in HTA assessments

Cases the HTA assessments considered input from:

internationally to enable informed decisions

about care management strategies

clinical experts people with MS



Specific aspects people with MS raised as not being adequately addressed

balance and dizziness



fatigue and weakness

Engage healthcare systems

Widespread adoption of current guidelines would strengthen diagnosis, treatment and monitoring

Incentives should support improvement in clinical practice and the incorporation

bladder problems

of new evidence on MS management in

healthcare decision making