SHORT REPORT

Forced hyperphasia and environmental dependency syndrome

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Abstract

A distinctive, language related fragment of the environmental dependency syndrome is described: compulsive, involuntary, environmentally dependent speaking. Because this syndrome represents the opposite of aphasia, it is named forced hyperphasia. An 84 year old woman with acute left frontal infarction was admitted to hospital with gait disturbance, forced grasp reflex, and striking imitation behaviour. After 2 weeks her imitation behaviour disappeared, but an equally striking new behaviour emerged. In the presence of others she would call out the names of objects in the room, and also call out the actions and gestures of people in the room, even though she was not asked to do so, and even though she was asked to stop. For example, if the doctor scratched his nose, she said, “The doctor is scratching his nose.” Brain CT, MRI, and SPECT showed cerebral atrophy and a left superior frontal subcortical infarct. It is suggested that “forced hyperphasia” is a clinical fragment of the environmental dependency syndrome and that her compulsive, impulsive, involuntary release of spoken language resulted from the release of frontal inhibition of the complex reflex linking environmental cues to the set of motor, limbic, spatial, and linguistic associations underlying spoken language.

Case report

An 84 year old, right handed woman was admitted to the hospital for evaluation of memory impairment and gait disturbance. One month before admission she had sudden onset of right sided weakness. After that episode she developed increasing complaints of memory impairment and gait disturbance.

Positive neurological findings on admission included bilateral grasp reflexes with forced visual groping with the right hand (visual grasp reflex). She also had other evidence of frontal system dysfunction, including snout, suck, and palomental reflexes. She had a mild right hemiparesis; she could not walk unaided; deep tendon reflexes were increased bilaterally; Babinski’s sign was present on the right. Pain sensation was decreased on the right.

Orientation to time, place, and person was within normal limits. Speech output was fluent with phonemic paraphasias. Repetition was limited to short sentences. She could not follow simple spoken commands, such as “open your mouth.” She had recurrent perseveration. She could read and carry out written commands. She was agraphic for Kana and Kanji. Writing to dictation for Kana letters was 80% accurate; for Kanji letters, 40%; and for simple sentences, 80%. Written picture descriptions were 50% accurate. Writing, both to dictation and on written picture description, was large,
subcortical infarction. FLAIR (fluid attenuated inversion recovery) image of MRI demonstrating the left frontal temporal areas.

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When asked why she did this, she said she felt the doctor touching his hair. If an examiner waved his hand, she said, “The doctor is waving his hand.” This “speaking out” behaviour was impulsive, compulsive, involuntary, and apparently, unstoppable. She never repeated the sentences of the examiner unless so requested. By the 30th hospital day, her forced hyperphasia disappeared.

Discussion

We think that forced hyperphasia is a language linked fragment of the environmental dependency syndrome. Patients with frontal lobe damage have been described as being “pulled to stimulus”\(^1\).\(^4\)\(^5\) Imitation behaviour and utilisation behaviour seem to be complex forms of “pull to stimulus.” Forced hyperphasia, first described to our knowledge by Shimomura et al.,\(^1\) probably falls within the same neurobehavioural family.

“Pull to stimulus,” of course, is simply another descriptive term for a behaviour which has been difficult to explain. The best current explanation, in our opinion, is that this behaviour is a complex reflex resulting from release of frontal inhibition. Among the functions of the frontal lobes is inhibition of inappropriate motor, limbic, and spatial responses which might impulsively be released as a reaction to disturbing or irrelevant stimuli. Such behaviour, a “pull to environmental stimulus,” is universal in infants. As the frontal lobes mature, instinctive, involuntary, reflex “pull to stimulus” is inhibited; the sensorimotor and association systems develop ever more complex, controlled interactions with the environment. A neurally based “social grammar” is developed whereby cognitive and emotional systems are integrated with the environment, and we develop the capacity to control our environment and plan ahead. The “social grammar” is the internalised, neurally conditioned set of rules which order and constrain interactions between and among people and their environment(s). Impulsive, inappropriate, environmentally linked, complex reflex responses are released when frontal inhibitory systems are damaged.

Why don’t we see environmental dependency syndromes (forced hyperphasia, forced hypergraphia, visual grasp reflex, imitation behaviour, utilisation behaviour) more often? Most likely the “social grammar”—that is, the rules whereby the brain interacts with the envi-
environment is multilayered in interlocking neural nets, widely distributed throughout the brain. Focal damage to the frontal lobes without more widespread damage to the neural substrate underlying the “social grammar” may not be sufficient to cause an environmental dependency syndrome. Lhermitte made the point that “loss of intellectual control” was required for the occurrence of such behaviour. Our patient had not only the frontal lobe lesion but also damage in other portions of the brain. Shimomura et al. described what we are calling forced hyperphasia in patients with dementia of the frontal lobe type. Exploration of the neurobiology of rules whereby the brain interacts with the environment (the neurology of the “social grammar”) should be a fruitful endeavour.

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