Amyotrophic lateral sclerosis (ALS) can result in the locked-in state (LIS), characterized by paralysis, and eventual respiratory failure, compensated by artificial ventilation,1 or the completely LIS (CLIS), with additional total paralysis of eye muscles. Brain–computer interfaces (BCIs) have been used to allow paralyzed people to regain basic communication,2 although current EEG-based BCIs have not succeeded with CLIS patients.3 We present Class IV case evidence to establish that communication in the CLIS is possible with a metabolic BCI based on near-infrared spectroscopy (NIRS).

Case report. In August 2012, a 67-year-old woman (video 1 on the Neurology® Web site at Neurology.org) was admitted to CERES, an institution for the care of apalic patients affiliated with Tübingen University, over a period of 4 weeks. The patient had been diagnosed with bulbar sporadic ALS in May 2007, as locked-in in 2009, and as completely locked in May 2010, based on the diagnosis of experienced neurologists. Her husband reported that she had lost the ability to communicate, which was last possible by using her eyes, approximately 27 months prior to admission (table e-1; e-Patient). She had been artificially ventilated since September 2007, fed through a percutaneous endoscopic gastrostomy tube since October 2007, and was in home care. No communication with eye movements, other muscles, or assistive communication devices was possible at the time of admission (score 0 of 48 on the ALS Functional Rating Scale–Revised)4 (video 2). Her cognitive functions were assessed by an extensive neurophysiologic examination based on event-related brain potentials (e-Assessment; figure e-1), suggesting intact cognitive functioning. Before this study, training using a similar paradigm with an EEG-based BCI did not result in stable successful communication.5 A second 2-week admission (second period) and a third 4-week period at home took place in 2013. The study was approved by the ethics committee of the Medical Faculty of Tübingen University and informed consent was obtained from the legal guardian.

Methods. Functional activations of the cerebral cortex to auditory processing of correct or incorrect statements were assessed with NIRS, which can be applied at the bedside, is comparatively low cost, and detects slow changes in hemodynamic brain responses (figure 1A). We employed blocks containing 10–20 auditorily presented sentences (factually true or false), with intersentence intervals (ISI) of 25 seconds, needed for the corresponding hemodynamic response to develop. There were sentences with known answers (e.g., “You were born in Hamburg”), and sentences with unknown answers, termed open sentences (e.g., “You want to be moved from left to right”). The patient was instructed to think “yes” or “no” after each sentence. Sentences were recorded with the voice of the husband (the primary caregiver), but in some sessions a person unknown to the patient spelled the questions to control for any bias. Feedback was given after the ISI: “Your answer was recognized as Yes,” or “Your answer was recognized as No” (figure 1B) depending on online classification (videos 2–4). We preprocessed the time-series data of the measured changes, extracted segments corresponding to the ISI of 25 seconds for each sentence (data point), and employed binary classification of yes vs no brain responses using support vector machines (e-SVM).

Results. Classification of the patient’s NIRS data from the 25-second period after sentence presentation, when the patient had to think the answer, yielded significantly different deoxygenation levels for the yes and no answers across all 3 training periods (figure e-2). The percentage of correct answers was 71.67% (t11 = 4.73), 75.71% (t27 = 7.38), and 76.30% (t26 = 8.43) for the yes and no answers across the 3 periods, all significantly different from chance, p < 0.01 (figure 1, C–E, e-Table e-2). The positive predictive value was 80.9% and the negative predictive value was 72.9%. The classification of the sentence-induced change in deoxygenation prior to the ISI yielded only chance-level results, suggesting that brain changes related to sentence presentation alone did not create the yes/no differentiation. Sessions 8 and 12 of the third period were separated by 11 days and contained the same questions with known answers. Both sessions resulted in 100% correct classification. Sessions 28 and 29, also separated by 11 days, contained the same open questions and resulted in 100% correct answers (e-Replication).

Discussion. Using NIRS, we obtained significantly above chance-level answers in a CLIS patient over an...
extended period of time. The overall performance of 76.30% in the last training period and the 100% correct performance in some sessions suggest that this is a viable method to reestablish communication in CLIS patients with a high test-retest stability. To generalize from a single case to all CLIS patients is not possible, but the accessibility and simplicity of NIRS methodology and the use of standard free access algorithms such as SVM should encourage and facilitate exact replications. Metabolic BCIs might thus break the unbearable “silence” of CLIS.

From the Institute of Medical Psychology and Behavioral Neurobiology (G.G.-A., A.F., K.T., C.A.R., N.B.) and Graduate School of Neural and Behavioral Sciences (G.G.-A.), University of Tübingen, Germany; Escuela Superior Politécnica del Litoral (ESPOL) (G.G.-A.), Ecuador; National Rehabilitation Center for Persons with Disabilities (K.T.), Tokorozawa, Japan; Central Institute of Mental Health (H.F.), Medical Faculty Mannheim, Heidelberg University, Mannheim, Germany, and Ospedale San Camillo (N.B.), IRCCS, Venice, Italy. Author contributions: G. Gallegos-Ayala and N. Birbaumer designed the brain–computer interface and performed the training. A. Furdea wrote the program for the assessment and analysis. K. Takano contributed to training, assessment, and analysis in the second part of the study. C.A. Ruf organized the training sessions and edited the report. G. Gallegos-Ayala, H. Flor, and N. Birbaumer wrote the report.

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ACUTE REFRACTORY INTESTINAL PSEUDO-OBSTRUCTION IN MELAS: EFFICACY OF PRUCALOPRIDE

In mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes (MELAS), a multisystem mitochondrial disorder, gastrointestinal involvement is frequent with dysphagia, chronic diarrhea, anorexia, abdominal pain, delayed gastric emptying, and paralytic, often intractable, ileus. In this article, we report a patient with chronic gastrointestinal dysmotility and acute refractory intestinal pseudo-obstruction responsive to prucalopride.

Level of evidence. This single observational study without controls provides Class IV evidence.

Case report. A 32-year-old man with typical MELAS associated with the A3243G mutation of mitochondrial DNA was admitted to our hospital because of the sudden onset of abdominal pain, nausea, vomiting, and failure to pass flatus or stool. He had a long-lasting history of gastrointestinal dysmotility problems only partially responsive to common prokinetic therapy. He was cachectic with prominent loss of skeletal muscle mass and a body mass index (BMI) of 17.3. Brain MRI showed the sequelae of previous stroke-like episodes but was negative for recent events. Abdominal examination revealed severe distention, tympani to percussion, and absent peristalsis. Extensive blood tests showed leukocytosis and lactic acidosis (serum lactate 13 mEq/L; normal range 0.5–22). After 2 days, the patient developed hemodynamic instability with a systolic blood pressure of less than 80 mm Hg and clinical signs of tissue hypoperfusion, low consciousness, oliguria, and tachycardia. CT scan showed diffuse dilation of the large bowel, air fluid levels, and abundant intraluminal fecal material with no signs of definite obstruction or bowel wall thickening (figure, A and C). Search for Clostridium difficile toxin A/B, botulinum toxin, and stoll cultures were negative. The patient, diagnosed with intestinal pseudo-obstruction, was put on total parenteral nutrition. Attempts with decompressive colonoscopy and therapy with IV metoclopramide (10 mg TID), oral erythromycin (250 mg BID), and IM pyridostigmine bromure (60 mg TID) for 1 week failed to produce any clinical benefit. Oral prucalopride (4 mg BID) was then added with rapid clinical improvement in 1 day and complete resolution of the colonic pseudo-obstruction in 10 days. Lactic acid decreased to 3.5 mEq/L. After 1 year of follow-up, the patient is still on therapy with oral prucalopride (4 mg daily) and has not experienced any further gastrointestinal symptoms. Moreover, he exhibits a marked improvement of fatigue and exercise tolerance. The current BMI is 21.5 and CT scan shows no pathologic findings (figure, B and D).

Discussion. Acute intestinal pseudo-obstruction is a clinical condition characterized by abdominal distention, nausea, vomiting, and abdominal pain. Radiologic features are dilation of the transverse and ascending colon simulating mechanical obstruction. Gastrointestinal dysmotility may represent a severe complication in mitochondrial encephalomyopathy, most commonly observed in mitochondrial neurogastrointestinal encephalomyopathy (MNGIE), but also in MELAS. The mechanism of paralytic ileus in MELAS is not completely understood and either a myenteric plexus neuropathy or a visceral myopathy has been hypothesized. Supporting the hypothesis of the visceral myopathy, a diffuse and profound cytochrome c oxidase deficiency has been recently demonstrated in the intestinal smooth muscle cells in all regions of the gastrointestinal tract. Thus, pathophysiology of intestinal pseudo-obstruction in these patients might be related to mitochondrial dysfunction and energy metabolism imbalance similar to what happens in the brain in stroke-like episodes. According to this, in our patient blood lactic acid was markedly elevated in the acute phase of paralytic ileus and returned to the pre-crisis level with clinical improvement. Paralytic ileus has been considered as an almost untreatable complication...
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Guillermo Gallegos-Ayala, Adrian Furdea, Kouji Takano, et al.
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