Pre-Syncope and Ataxia After Spinning Whiplash

Abstract
This is a 46-year old female presenting to her Primary Care Physician on Day #2 post-motor vehicle accident. She was the driver of a large suburban truck that was suddenly T-boned into her left drivers’ rear wheel, whirling 360 degrees to face oncoming traffic. She and her daughter went home. But over time, the patient slept more and more. She experienced ataxia and pre-syncope. Dozens of presentations to the Emergency Department (ED) ensued. Eventually, she could not stand up without vertigo, ataxia, pre-syncope, and vomiting. After 2 months, the patient still had severe nausea and vomiting 12 times/day, was weak, ataxic, confused, and had short-term memory loss, and developed tinnitus and an expressive aphasia. She still had pre-syncope on standing. She was diagnosed with a traumatic brain injury (TBI).

After 3 months, the patient underwent an MRI of head and neck without contrast, and it was negative. Ten ED visits later, the doctor ordered a spiral CAT scan that detected an extra cranial 3 cm left vertebral artery dissection (VAD). Angiogram confirmed the VAD at C4-5, with a 2 cm aneurysm at the base. Medical management failed to correct pre-syncope, so the patient kept “doctor shopping” until led to Electrophysiology Cardiologist, Dr. David Cannom, MD. She underwent the Tilt Table Test (TTT), passing out for 22 seconds. The diagnosis of Dysautonomia was made. This was aggravated by polydypsia and polyuria, so she was diagnosed with diabetes insipidus (DI) and placed on a PICC line for almost four years.

After 7 years and a nearly full recovery with a return to work as an anesthesiologist in sight, a neurologist allowed her fall to the hard wood floor during his “close your eyes and touch your nose” exam while standing; the patient sustained another TBI with DI and remains disabled.

Keywords: Pre-syncope; Vomiting, Ataxia; Traumatic brain injury; Diabetes insipidus, Vertebral artery dissection; Dysautonomia; Oculo-vestibular dysfunction; Gastroparesis; Syncope

Abbreviations: DI: Diabetes Insipidus; ED: Emergency Department; EPS: Electrophysiology; ICU: Intensive Care Unit; IV: Intravenous; PICC: Percutaneous Intravenous Central Catheter; POTS: Postural Orthostatic Tachycardia Syndrome; TTT: Tilt Table Test; TBI: Traumatic Brain Injury; VAD: Vertebral Artery Dissection

Introduction
This case highlights the issues of “physician patients,” “self-diagnosis,” “malingering,” dysautonomia, TBI with DI, and VAD. The question of whether some illnesses are “rare” vs “undiagnosed” is a key focus, as well as the proposed need for increased physician education on “invisible illnesses,” so that patients are not distraught and made to feel mentally ill. The sequelae of depression, anxiety, and loss of patient dignity contribute greatly to decreased quality of life, and increased monetary strain on the health care system, which in some cases, is being used out of desperation, as a “revolving door.” Failure to diagnose could contribute to physician frustration and loss of job enthusiasm.

Case Presentation
While still a resident in Anesthesiology at the University of Southern California School of Medicine, the patient initiated the grant-writing and research of patients with intractable closed head injury in the Intensive Care Unit (ICU). It was published as a Rapid Communication in Neurosurgery [1]. She continued academic medicine pursuits in the subjects including the Surgical ICU [2-5], pulmonary hypertension [6], nitric oxide [7,8], ventilation/perfusion mismatch [9,10], and Hyperpolarized 3He-MRI of the porcine lung [11-13]. She also presented multiple Poster Presentations and was a leader in her field, Hyperpolarized 3He-MRI of the porcine lung, having internationally lectured on V/Q mismatch and 3He-MRI of the lungs [14,15]. She collaborated with other anesthesiologists, radiologists, internal medicine physicians, and physicists on projects [11-13] and helped write $3 million in successful NIH Research Grants at the University of Pennsylvania.

Being keenly astute as to her symptoms and medical history, she was unprepared to undergo the path to her multiple “rare” diagnoses, yet was known for her “persevering” attitude. Prior to the car accident, her last job position was as anesthesiologist and Director of the SICU at a Veterans Administration in Southern California. Her highest position was that as Interim Chief of Anesthesiology at the Philadelphia Veterans Administration in Pennsylvania, USA. She had no medical problems except seasonal allergic asthma, anaphylaxis to sun exposure, and anxiety due
to job-related stress and a pregnancy that was carried to term, with elective Caesarian Section and no complications except for mastitis, for which she was treated with antibiotics.

This is to be clear that she was not only an intelligent physician, but an academic one who was used to defending her research by symposium podium or poster presentation, or lecture hall podium in Grand Rounds. She also had expertise in making quick and authoritative decisions both as Chief of Anesthesiology and in the Operating Rooms, as an anesthesiologist, and as an intensivist trained in the ICUs at Stanford. She had a noted reputation for being extremely ethical, compassionate to all of her patients, and for never “losing her cool.”

Discussion

TBI with DI

The CDC estimates 1.7 million TBIs per year, with about 30% leading to death. This combination is associated and learned by every medical student, with there being no need for actual head impact for TBI to occur, as in whiplash. The patient presented with poor cognition, inability to drive without getting lost, forgetfulness, and an expressive aphasia. Physical exam findings included nystagmus and loss of proprioception when standing alone. This was also associated with nausea/vomiting, diagnosed as mal de Debarquement Syndrome. Both CT scan and MRIs were normal for brain anatomy.

DI

The incidence of severe DI in acute TBI is about 3%. Secondary hypoadrenalism and hypothyroidism can occur in a secondary or later stage of TBI, with an incidence of about 10%. There is a need to evaluate the pituitary: hypothalamic axis, the thyroid axis, and ADH secretion.

Each time, the DI was diagnosed by the patient. Polyuria and polydipsia were the primary presentations. With central or psychogenic DI, the posterior pituitary gland is traumatized, leading to a defect in the hypothalamic: pituitary gland axis, which regulates ADH and total body water. No ADH is produced. The diagnosis can be made by the water deprivation test, if the patient is not imminently close to renal failure from dehydration. This involves [1] maintaining npo status, [2] insertion of a Foley catheter, and [3] water deprivation. In the first DI episode, the patient urinated 4 liters urine in 2 hours. On the second episode, she was dehydrated by lab results: BUN/CR =108/4.1; GFR = 35%; platelet count of 45K on generic clopidogrel 75 mg (for the VAD) and simply treated with IM DDAVP and resuscitation prior to renal insult. Resuscitation included NS fluid bolus plus 60 ml/ hr; IV steroids, meclizine, and head down flat. Legs were elevated for a mean arterial pressure less than 60 mm Hg.

Treatment of DI was with anti-diuretic hormone (ADH) po, AKA DDAVP via nasal spray. The underlying dysautonomia caused titration problems so that sometimes the see-saw tipped her sodium to 158 mcg/ml and she was dehydrated; other times, her sodium was 127 mcg/ml and she felt like a water balloon about to undergo uncal herniation and eyeball expulsion. Both nasal and po DDAVP were problematic as treatment therapies, due to the underlying dysautonomia as a hypovolemic state. The patient relied on daily weights and the color of her urine (clear water vs. dark yellow) to dictate the next dose of DDAVP, especially overnight. Her chief complaint was dry mouth, which required more frequent dental visits to avoid periodontitis due to lack of sterilizing saliva. This put her at risk of cerebral stroke, especially since laminar flow in the vertebral artery dissection area could not be guaranteed, tending towards coagulation. Each time, the DI resolved after about 6 months.

Dysautonomia

The incidence of orthostatic hypotension is age-dependent, with patients under 50 yo having a 5% incidence, and patients over 70 yo having a 30% incidence. The patient experiences significant quality of life decreases, with increased mortality and incidence in heart attack, stroke, heart failure, and atrial fibrillation [16].

In the ED, patient presentations are wrought with disgust and dread. Doctor after doctor did orthostatics on the patient, with her standing numbers going as low as a blood pressure of 75/44 mmHg, with a heart rate of 110 bpm. When she was unable to stand up for fear of syncope, one doctor said he was “going to look it up in my medical school books” and came in later to say, “You’re supposed to wait 5 minutes before taking the standing blood pressure.” The patient said, “Do you want to scrape me off the floor? I won’t last that long!” There was a large dichotomy between physician assessment/impression and patient complaints/experience [17-19]. Only because one doctor knew another doctor who “worked with patients who faint,” did the patient finally get a referral to an EPS cardiologist. The most common presentation of orthostatic intolerance is in a female (80%) with Postural Orthostatic Tachycardia Syndrome (POTS), which is defined as sustained increase in heart rate by 30 bpm at 10 minutes after standing, with an increase in noradrenaline serum concentration [20].

The patient was checked in to the Outpatient Cardiology Suite for the state-of-the-art diagnostic tool, the Tilt Table Test. She complained of “going gray” when the table was lifted from the supine position to a 90° standing angle. She had complete syncope with loss of consciousness for 22 seconds, without seizure activity or loss of bowel/bladder function. The definitive diagnosis of Dysautonomia was made. The pathophysiology of POTS and other dysautonomias is an underlying cerebral, brainstem hypoperfusion upon standing, with lack of vasoconstrictor alpha-1 agonist activity in the legs.

Since then, she has been on the alpha-agonist Midodrine to increase her blood pressure, she wears Jobst® Stocking thigh-highs, and an abdominal binder. All of these increase cerebral blood flow and allow perfusion to the brainstem primary autonomic nervous system centers, allowing her to sit and stand for small periods of time. She has spent most of the last 9 years in bed.

Only in the cardiac hospital, Good Samaritan, where Dr. David Cannom practices with many patients with Dysautonomia, was she ‘fast-tracked’ into the ER, into admissions, into getting an iv consults, and getting her brain re-perfused with bolus NS, iv steroids, meclizine, ondansatron, and dilaudid as “her” recipe back to health. After 4 years, the PICC line was removed. But the

dysautonomia kept her in a constant hypovolemic state by nature, and she was taking 10 mg Midodrine every morning, and another 10 mg in the afternoons. For nausea/vomiting, she was on IV Ondansetron 4 mg bid for 3 years, then on 4 mg/day until now. Endocrine workup showed a diagnosis of hypothyroidism, which was treated with ½ gr Armour Thyroid.

Even after the diagnosis of Dysautonomia was made, not one doctor had ever heard of “Dysautonomia,” so the reactions were genuinely a social study to observe. Some doctors pretended to know what it was, whereas others just looked blankly at the patient. Only a few asked for an explanation. If an unsolicited explanation was offered to benefit patient care, all smiles turned to frowns and the patient was treated as a “malingering.” Her requests for 1 liter of NS bolus were ignored, as were her requests for treatment of a migraine headache. She had to leave the ED by wheelchair without any treatment on several occasions for lack of any treatment at all. She successfully argued against paying the ED insurance premium for lack of treatment. All in all, though, it was humiliating.

Vertebral artery dissection with aneurysm

Extracranial vertebral artery aneurysms are rare and not well described. For a small subset of patients with underlying hereditary or connective tissue disorder, operative intervention may be a choice through exclusion and reconstruction. The author surmises that many VADs simply go undiagnosed due to patient death.

Once the spiral CT detected a left vertebral artery abnormality, the patient was sent by ambulance to a major medical center for an angiogram of the brain. Upon angiogram, she recalls the injection and the warmth of the dye going through the Circle of Willis and back down the contralateral side. The extracranial IJVAD was further identified as being 3 cm long, with a 2 cm aneurysm at the base. Lots of discussion was held over treatment for VAD. Very little literature exists on this subject, and the multi-center studies still had a very small “n.” The final treatment plan was medical management without surgery or stenting: clopidogrel 75 mg/day for life.

The doctor was now the patient, and she was confused from Day #1. The ensuing events were remarkable not only for the fact that multiple, rare diagnoses were made, but that she had to “Speak up” and “Stand up” for her rights as a patient, refusing to be abandoned in a nursing home. She refused to be treated as a “drug-seeker.” She knew to ask for the Charge Nurse, and then she knew to ask for the Incident Report after the doctor allowed her to fall to the floor, sustaining not only the second TBI with DI, but also requiring Left Shoulder Rotator Cuff Repair for falling onto her left side and protecting her head from hitting the floor.

After 4 weeks of Neurorehabilitation and learning how to walk and talk again in 2006, the patient slowly recovered. In 2013, she had been accepted into Stanford’s Advanced Fellowship for Clinical Anesthesiology to “get my stethoscope back” but the 2nd TBI put this permanently out of reach.

It behooves physicians to listen to their patients, to take them seriously, and to refer them to another specialist (instead of sending them home empty-handed) if they are unable to make a diagnosis. Sometimes, physicians really need to “wrack their brains” hard, to find some solution for the patient. Some patients suffer for weeks, months, years, or decades until finally getting an “invisible illness” diagnosed by the 30th doctor! May this never be you, or one of your loved ones!

By increasing physician education on a handful of “invisible illnesses,” [20] many more diagnoses can be made in a timely manner, more referrals to the proper specialists can be made, and both patient and physician frustration should be replaced by the genuine joy and hope that may be otherwise lacking in this scenario. This testimony is an inspiration to both doctor and patient alike, for “Perseverance” trumps medical textbooks and medical Case Reports every time.

The patient is still disabled, has severe orthostatic hypotension, and some cognitive dysfunction is expected to remain. She has become a published author and avid Patient Advocate for “Invisible Illnesses,” serving on the Advisory Board to the Invisible Disabilities Association, which awarded her the 2011 Perseverance Award. She is a Public Figure on Facebook, and much loved by millions of disabled people who felt that doctors were simply “out of touch” and not sympathetic to the plight of those whose diagnosis may be best determined by another specialist. So, doctor, just because you haven’t heard of a disease, it doesn’t mean that it doesn’t exist. Stanford’s Medicine-X program encourages patients with invisible or rare diseases to join internet groups, become “e-patients,” then “e-scholars” who help other patients to self-diagnose a rare disease. So don’t squeak and squawk if your patient knows more about Chiari 1 malformation than you do. In today’s world, she has to know more in order to survive. Support her, as we are all working toward the same goal: health, longevity, and quality of life.

References


