Neurology outreach clinic for Huntington disease in Peru
Lessons for neurodegenerative diseases

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With increasing life expectancies worldwide, providing longitudinal care for patients with neurodegenerative diseases in low-resource settings is a growing challenge.1 According to the 2015 Global Burden of Disease Study, Alzheimer disease, other dementias, and Parkinson disease together affect more than 50 million people globally.2 Furthermore, the WHO estimates that at least 60% of these cases occur in low and middle-income countries.1 Although currently available treatments for neurodegenerative diseases are largely supportive rather than curative, they can reduce morbidity and burden of disease.1 As more effective medications become available and the prevalence of neurodegenerative disease increases, the need for skilled care providers will also increase.

Outreach clinics, which deliver specialty-level care to geographically remote areas, have been used for vaccination campaigns, HIV control, and obstetrics, as well as neurologic conditions like epilepsy and stroke. However, these clinics have not been used for the management of neurodegenerative diseases, which occur worldwide but at higher rates in populations bearing specific genetic risks. For example, Huntington disease (HD), a fatal hereditary neurodegenerative disease, is particularly prevalent around Lake Maracaibo in Venezuela and in the Cañete Valley of Peru.3 In this report, we present our experience with an outreach clinic focused on HD in Peru, with lessons that may be broadly applicable to neurodegenerative disease care in rural, low-resource settings.

The outreach neurology clinic program

The Neurogenetics Research Center (NRC) of the Instituto Nacional de Ciencias Neurológicas (INCN) developed an outreach clinic for the rural Cañete valley, where HD prevalence is 4-fold greater than the global rate.3 Cañete valley is only a 3-hour drive from the capital city of Lima; however, it has limited available medications, basic health infrastructure, and a single neurologist for more than 100,000 people. Many local patients with HD are unable to access care in Lima due to the expense of travel and mobility limitations, which worsen with disease progression. Notwithstanding the lack of medical personnel in Cañete, Peru is an upper-middle-income country and has multiple existing neurology residencies that make it possible to staff an outreach clinic.

The initial approach to the Cañete HD community was informed by an earlier incident that compromised trust in outsiders coming to focus on HD. In 2005, journalists, accompanied by some doctors and researchers, came to Cañete to interview and document the experience of HD families. Patients and their families, including children, were photographed without obtaining appropriate consent or informing the families what would be done with the photographs and information. The resulting journalistic piece was a sensationalistic and stigmatizing piece called “The Curse of Cañete,”4 which described how whole families are affected by...
HD, how it occurs at a high prevalence in people from Cañete (implying that contact with people from the area could bring about the curse), and quoted a doctor saying that these people should have never married. It also used original patient and family member names and displayed unflattering photographs. The piece ran in Cañete as well as Lima, and many of the HD patients and families felt betrayed by the journalists as well as the researchers, doctors, and community agents that had facilitated the journalists’ access. This history led the NRC team to prioritize community buy-in and engage a local community agent for outreach, institute strict consent processes for genetic testing as well as any research protocols, limit photography, and be more judicious in bringing outside visitors to the outreach clinics.

The outreach clinic consists of 2-day visits to Cañete every 3 months, which have been conducted since 2014 by a multidisciplinary group of health care providers (neurologists, psychiatrists, geneticists, residents, students, and support personnel) traveling from Lima. They divide into 2 teams, with one attending to patients at a local community health center, and the other conducting house visits for homebound patients throughout the region (figure). During the trips, the outreach clinic staff provides neurologic and psychiatric consultations and follow-up visits, distributes medication brought from Lima, takes samples for genetic testing, and provides counseling and family support. Patients (or occasionally their family members) come back to each outreach clinic for medication refills and follow-up, and there are some new patients, as well as some deaths and losses to follow-up. Clinic personnel conduct HD-related training for caregivers and local health care providers and recruit participants for ongoing research. Although the clinic focuses on HD, about half of the patients have other neurodegenerative conditions such as Parkinson disease, ataxia, or dementia (figure).

This program is financially supported by local public–private collaboration. The public component is funded through resources provided by the INCN (van transportation, laboratory supplies, excused hours for contracted members of the NRC team), the Cañete regional health system (meals and lodging), and the government health insurance, Seguro Integral de Salud (SIS) (provides medicines for almost half of the patients). The private component consists of medication donations from local pharmaceutical dispensaries for patients who do not have SIS, as well as volunteer time and funds from the NRC team (other meals, office supplies). Official summaries of the clinic activities, services, and finances are generated for the regional government within a few weeks of each clinic’s completion. The data in the figure were generated from 7 summary reports for the 18-month period from March 2015 to October 2016.

Lessons from the outreach clinic experience

The outreach clinic provides many lessons that may be applicable for similar chronic conditions:

1. Neurodegenerative disease care requires consistent follow-up (table, excerpt 1). Many neurologic drugs require

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**Figure** Clinic resources, structure, and services

*Medications typically include antipsychotics (risperidone, olanzapine, haloperidol), selective serotonin reuptake inhibitors (citalopram, sertraline), benzodiazepines (clonazepam), carbidopa-levodopa, valproate, topiramate, and others. **Parkinsonism, dementia, ataxias, headache, epilepsy, neurodevelopmental disorders. HD = Huntington disease.
Table: Huntington disease (HD) patient vignettes and caregivers’ perspective on outreach clinic

**Patient vignettes**

A 31-year-old woman with aggressive HD psychosis, locked in her room for safety by family. Able to interact with family and come out of the house after 9 months of treatment with second-generation antipsychotics and psychiatric consultations.

A 42-year-old man with HD, moderate chorea, falls, and debilitating depression. Able to return to work on his farm and maintain status as breadwinner for family with young children after 6 months of treatment with second-generation antipsychotics and antidepressants.

A 4-year-old boy with juvenile HD, severe developmental delay, ataxia, and constant, daily seizures. With antiepileptic medications and frequent follow-up, seizures were controlled for 6 months.

**Topic** | **Interview excerpts**
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1. Importance of consistent access | Caregiver: I have to wait 2 months for when you come with the campaign so you can see him. But in those 2 months, things can happen and I have to wait. As I’ve said, the disease, it’s not that you come with the campaign and that’s it, and you are good, no, the disease is constant... We should have... everyone who has a family member with this condition, someone specialized 24 hours per day.
2. Preference for home visits | Interviewer: Would you prefer to have doctor’s visits here in your home or in the health center?
Caregiver: It’s better that they come to my house.
3. Role of social worker | Caregiver: Today I think the (social worker) from the health center sent me a message to bring him. I communicated this to his brothers and he said, ok, bring him.
4. Appreciation for outreach clinic | Caregiver: At least now there are more campaigns, no? For example they come to the house, visit him, give him medicine, and do the evaluation, no? Of every medicine, to see if he has progressed or has not progressed, no? In this respect it’s good what they are doing now.
5. Financial difficulties | Caregiver: Since in the farm we do not have money available... so it costs for a medical visit, it costs too much. And the pills are very expensive. These tablets... we earn 30, 40 soles in the field... it’s very hard.

* Formal qualitative analysis, with thematic saturation of outreach clinic-related topics in interviews, coding, and theme development, were not conducted; topics represent common sentiments expressed via personal communications and in response to exploratory questions about the outreach clinic.

Quotations were originally acquired for a larger study on quality of life in Huntington disease, which received institutional review board approval from the Instituto Nacional de Ciencias Neurológicas and required written informed consent from all participants.

Challenges and opportunities for the future

1. Defining measurable outcomes for evaluating the efficacy of an intervention in neurodegenerative disease care. Despite the documented qualitative effect of the program on patients and caregivers (table, excerpt 4), the phenotypic heterogeneity of neurodegenerative conditions makes it difficult to establish precise and objective outcome measures for the program. In HD, for example, clinical improvement due to proper management can manifest in a myriad of ways that are hard to quantify: extra years of employment, fewer falls, control of behavioral disturbances, or decreased caregiver burden. HD-specific health-related quality of life questionnaires are currently being validated in the Peruvian population and hold promise for future outcome evaluations.

2. Developing local health capacity for neurodegenerative disease care and expanding interdisciplinary involvement. We have been holding regular trainings and lectures with local health care providers, but it has not yet been feasible...
for them to make medical decisions regarding HD care. In addition, although we have genetics, psychiatry, neurology, nursing, and social work on our team, we have yet to involve nutritionists, geriatricians, or physical therapists in the outreach clinic.

3. Obtaining sustainable funding. The public–private outreach clinic funding model allows for integration into the existing Peruvian health system and provision of care for the poorest patients (table, excerpt 5). However, significant bureaucracy and personnel changes in government sometimes generate unpredictability in funding. International funding and more extensive collaboration with local nongovernmental organizations would provide more stability and allow the clinics to grow.

Although HD and other neurodegenerative diseases are not typically considered a priority in low-resource settings, their effect on patients, families, and the surrounding community can be devastating. The strategies used in this Peruvian neurology outreach clinic model can be replicated to provide much-needed specialty care for patients with neurodegenerative disorders in low-resource and remote settings throughout the globe.

Author contributions
Anastasia Vishnevetsky: study concept and design, analysis and interpretation of data, critical revision of manuscript for intellectual content. Maryenela Illanes Manrique: acquisition of data, critical revision of manuscript for intellectual content. Miguel Inca-Martinez: acquisition of data, critical revision of manuscript for intellectual content. Karina Milla-Neyra: acquisition of data, critical revision of manuscript for intellectual content. Elison Sarapura-Castro: analysis and interpretation of data, critical revision of manuscript for intellectual content. Pilar Mazzetti-Soler: study supervision, critical revision of manuscript for intellectual content. Mario Cornejo-Olivas: study concept and design, critical revision of manuscript for intellectual content.

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Disclosure
The authors report no disclosures relevant to the manuscript. Go to Neurology.org/N for full disclosures.

References

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