

# GAD65 and Glycine Receptor-Associated Neurologic Autoimmunity and Stiff-Person Syndrome within the University of Utah Health Care System

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# Objective

Describe epidemiological characteristics, antibody status, and treatment outcomes of stiff person syndrome patients within University of Utah Health

# Background

Stiff person syndrome (SPS) is an autoimmune disease that classically causes co-activation of agonist and antagonist muscle groups leading to severe muscle rigidity and spasms.<sup>1</sup> GAD65 is the most common associated neuronal autoantibody; patients may also have associated glycine receptor (GlyR) or amphiphysin autoantibodies.<sup>2</sup>

The incidence of SPS is estimated at 1 to 2 per million people.<sup>3</sup> Average age of onset is 40 years and females are more commonly affected.<sup>4,5</sup>

Diazepam and baclofen are the primary symptomatic treatments. Intravenous immune globulin (IVIg) is often an effective immunotherapy.<sup>5</sup> Large randomized, controlled trials and treatment guidelines for immunotherapy are lacking.

# Methods

Retrospective review of patients within the University of Utah Healthcare system from 7/1/2010 to 11/7/2018, meeting criteria 1 and 2, or 2 and 3, and crossreferenced for internal consistency:

- 1.Positive GAD65 (>100 IU/mL) or amphiphysin serum or cerebrospinal fluid testing at Associated Regional and University Pathologists, Inc. (ARUP Laboratories), or positive GlyR testing (performed on a research-basis) at Mayo Medical Laboratories.
- 2.At least one visit with a University of Utah clinician in the Department of Neurology.
- 3.At least one ICD code for stiff person syndrome (333.91 or G25.82), encephalomyelitis (323.9 or G04.90); or ICD code for GAD65 seropositivity (R76.0)

Patient charts were reviewed and further epidemiological and treatment data was collected on patients who had clinical symptoms consistent with SPS or variants along with associated seropositivity or additional clinical data supporting a SPS diagnosis.

# Acknowledgements

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## **Epidemiological Data**

Total # of SPS patients

Male

Female

Alive

Deceased

Caucasian

Other (Sudanese)

Mean reported age of diagnosis

Mean age at diagnosis

### Other associated antibodies n = 18

ANA

TPO and/or thyroglobulin

Intrinsic factor or gastric parietal Other

#### Visual symptoms

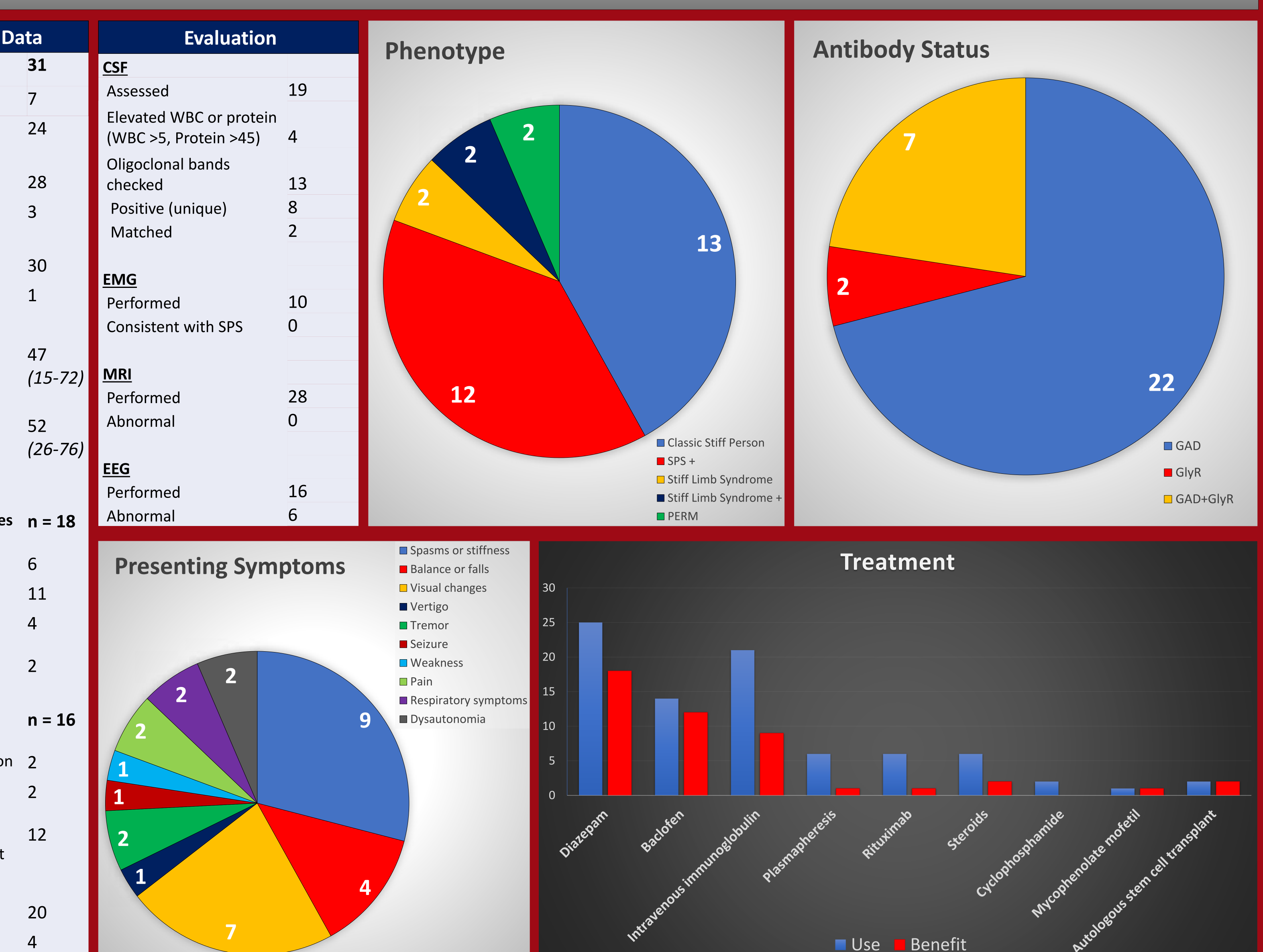
Positive visual phenomenon 2

- Visual acuity changes or scotoma
- Diplopia, nystagmus, or abnormal eye movement

Associated Autoimmunity Associated Malignancy

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# Results





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# **Discussion and Conclusions**

We identified 31 patients with SPS within the University of Utah healthcare system. Patients were predominantly female (78%) with the most common phenotype being classic SPS or SPS with concomitant cerebellar symptoms or epilepsy.

Our cohort had co-existing autoimmune disease (63%) and malignancy (13%).

GAD65 was the most common associated antibody; GlyR antibody testing should be considered in patients with SPS phenotype with negative or low GAD65 titers.

Diazepam and baclofen was effective in a majority of patients. IVIg was the most commonly utilized immunotherapy (used in 69% of patients) with benefit demonstrated in 41% of patients who received this treatment. Other immunotherapies demonstrated more limited benefit.

*Limitations:* Due to the electronic search method that was utilized, an accurate incidence could not be calculated. Treatment outcomes need more objective measures. The 2 patients who underwent bone marrow transplant were too early in their course to present additional data.

### References

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