Vagus nerve ultrasound in transthyretin familial amyloid polyneuropathy: A pilot study

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Abstract
Background and Purpose: Autonomic dysfunction is common in transthyretin familial amyloid polyneuropathy (TTR-FAP). Because ultrasonography is a powerful tool to study peripheral neuropathy, vagus nerve (VN) ultrasonography was used in our study to investigate the possible changes of the dimension of VN in TTR-FAP.

Methods: Eighteen patients with TTR-FAP and 17 age- and gender-matched individuals without any neuropathies were enrolled in a pilot study. The cross-sectional areas (CSAs) were measured bilaterally on transverse scans of vagus, median, and ulnar nerves. Clinical data were collected to explore the correlations with CSAs of VN.

Results: The median CSAs of VN in TTR-FAP were 3.5 (2.0-6.0) mm$^2$ on the right side and 2.5 (1.0-6.0) mm$^2$ on the left side, compared with 2.0 (1.0-3.0) mm$^2$ and 1.0 (1.0-2.0) mm$^2$ for healthy controls (HCs). There was a significant difference between the two groups on both sides ($p < .001$). The mean VN CSAs were correlated positively with the course of disease ($r = .7203, p = .0016$)(not including the patient with the longest disease course), the Composite Autonomic Symptom Score 31 ($r = .5252, p = .0252$), the left ventricular posterior wall thickness ($r = .5426, p = .0200$), and the interventricular septum thickness ($r = .5103, p = .0305$). The cutoff values of right and left VN CSAs to identify TTR-FAP from HCs were 2.5 and 1.5 mm$^2$ and the areas under the curve were .9395 and .8856, with a high sensitivity (.889 and .889) and specificity (.941 and .765), respectively.

Conclusion: VN enlargement is prevalent among TTR-FAP patients. VN ultrasonography may be an important clinical tool for assessing the severity of autonomic dysfunction in TTR-FAP.

KEYWORDS
CSA, transthyretin, TTR-FAP, ultrasonography, vagus nerve

INTRODUCTION
Transthyretin familial amyloid polyneuropathy (TTR-FAP) is a multiple systemic disorder caused by TTR gene mutation and characterized by deposition of transthyretin-derived amyloid fibrils in peripheral nerves, the heart, and other organs. Autonomic dysfunction is a common, early, and distressing condition of TTR-FAP that impacts quality of life, morbidity, disease progression, and mortality of patients. The autonomic neuropathies as initial symptoms struck 27.8% patients. Once symptoms were present, their evolution was equivalent to the...
progression of the motor and sensory neuropathy. Although the clinical variability of TTR-FAP has been well characterized as neuropathies, cardiomyopathy, or mixed phenotype, the extent of autonomic involvement remains poorly understood.

Detection of autonomic neuropathy is critical to identifying TTR-FAP from adult-onset neuropathies, such as chronic inflammatory demyelinating polyneuropathy (CIDP) and diabetic peripheral neuropathy (DPN). Because early differentiation of TTR-FAP from other peripheral neuropathies is of vital importance for early treatment, a number of studies related to detection of autonomic neuropathy have been performed for differential diagnosis, such as the eutectic mixture of local anesthetics test, laser Doppler flowmetry for vasomotor aspects, the Sudoscan and Neuropad test for sudomotor aspects, and skin biopsy. However, these approaches are invasive, complicated, or expensive, thus limiting their applications in clinical practice.

Nerve ultrasound is a painless tool for quick evaluation of peripheral nerve morphology. Previous ultrasound studies showed nerve enlargement in TTR-FAP, especially in the proximal segments of sensory and motor nerves. Vagus nerve (VN) ultrasound has been studied in DPN, CIDP, multifocal motor neuropathy, Charcot-Marie-Tooth Type 1A, Charcot-Marie-Tooth Type 1B, amyotrophic lateral sclerosis (ALS), and Parkinson’s disease. Abnormalities of VN are observed in all aforementioned diseases except ALS. VN, containing 80% of the unmyelinated fibers in the mid-cervical area, is considered to be related to autonomic dysfunction. Compared with other adult-onset neuropathies, autonomic dysfunction is more predominant in TTR-FAP. However, there has been no study on the VN ultrasound in TTR-FAP so far. This study was intended to find out whether VN ultrasound may be a useful tool for evaluation of disease progression and an early sign of autonomic neuropathy in TTR-FAP.

METHODS

Subjects

Between June 2020 and October 2021, 18 patients (14 males and 4 females) with TTR-FAP and 17 age- and gender-matched healthy controls (HCs) (13 males and 4 females) were recruited in Peking University First Hospital. All TTR-FAP patients were diagnosed according to the diagnostic criteria. The exclusion criteria of HCs were as follows: (1) skin numbness or paresthesia; (2) muscle atrophy or weakness; (3) other disorders of the peripheral nervous system; and (4) chronic diseases of other organs (eg, the heart, brain, eye, or kidney). The study was approved by the Ethics Committee of Peking University First Hospital. All participants provided written informed consent to participate in this study. All TTR-FAP patients with TTR gene mutations were inquired about their disease history and had a focused neurological examination and measurement scales performed, including the progressive course of disease, Coutinho stages (stage 0, no symptoms; stage I, unimpaired ambulation, mostly with mild sensory, motor, and autonomic neuropathy in the lower limbs; stage II, assistance for ambulation required, mostly with a moderate motor, sensory, and autonomic impairment of the four limbs; stage III,
wheelchair-bound or bedridden status with severe sensory, motor, and autonomic involvement of all limbs, Neuropathy Impairment Score (NIS), Norfolk Quality of Life-Diabetic Neuropathy Score (QOL-DN), and Composite Autonomic Symptom Score 31 (COMPASS 31). Cardiac parameters including serum N-terminal pro-brain natriuretic peptide (NT-pro BNP), interventricular septum thickness (IVST), and left ventricular posterior wall thickness (LVPWT) were collected.

**Ultrasonographic studies**

All subjects, whose bilateral VNs were measured and recorded by using the Philips Imaging System (iU Elite, Bothell, WA, USA), underwent peripheral nerve ultrasound. To be more specific, the 10-MHz high-frequency linear array probe was used. The cross-sectional areas (CSAs) of bilateral VNs were measured by tracing just inside the hyperechoic rim of the nerve. The transducer was positioned transversely in the lateral neck, and the nerve was identified inside the carotid sheath between the internal jugular vein and carotid artery (Figure 1). The variability from side to side was defined as the "maximum CSA/minimum CSA". The bilateral median and ulnar nerves were also measured at 10 sites according to our previously described method. The maximum CSA of median/ulnar nerves was defined as the maximum CSA of the 10 sites, and the mean CSA was the mean of the 10 sites.

Statistical analysis

IBM SPSS Statistics, version 24, was used for statistical analysis. The CSAs of HC and TTR-FAP showed an abnormal distribution (as evaluated by single sample K-S test). Thus, Mann-Whitney U test was used for evaluating differences in CSAs between TTR-FAP and HC. Receiver operating characteristic (ROC) curve analysis was performed to evaluate the applicability of CSAs measurements to differentiation of TTR-FAP from HC groups. The area under the curve (AUC) was calculated. The value of Youden index at its maximum was taken as the cut point for the diagnosis of TTR-FAP, and the sensitivity and specificity were calculated. Two-sided p-values were calculated for all analyses; p < .05 was considered significant. Spearman analysis was used to test the correlation between CSAs of VNs and other data.

**TABLE 1** Demographic and clinical characteristics of transthyretin familial amyloid polyneuropathy patients and healthy controls

|                    | TTR-FAP |             | p-value | HCs |             | p-value |
|--------------------|---------|-------------|---------|-----|-------------|---------|
|                    | Total, n = 18 | Early-onset, n = 9 | Late-onset, n = 9 | Early-onset vs. late-onset | n = 17 | Total TTR-FAP vs. HCs |
| Age at examination (years) | 50.4 ± 11.7 | 43.0 ± 12.4 | 57.8 ± 4.2 | .0035** | 47.6 ± 12.8 | .5182 |
| Male, n (%) | 14 (77.8) | 7 (77.8) | 7 (77.8) | >.9999 | 13 (76.5) | >.9999 |
| Positive family history, n, (%) | 13 (72.2) | 9 (100.0) | 4 (44.4) | .0294 | NA | NA |
| Coutinho stage I, n (%) | 10 (55.6) | 6 (66.7) | 4 (44.4) | .6372 | NA | NA |
| Duration of symptoms at examination, years, median (range) | 2.75 (0.7, 5.0) | 3.0 (0.7, 5) | 1.5 (0.8, 4.0) | .6173 | NA | NA |
| Height (m) | 1.7 ± 0.07 | 1.7 ± 0.07 | 1.7 ± 0.08 | >.9999 | 1.7 ± 0.7 | .2537 |
| Weight (Kg) | 63.6 ± 9.7 | 66.1 ± 11.3 | 61.1 ± 7.8 | .4201 | 66.5 ± 6.5 | .2090 |
| BMI (Kg/m²) | 21.7 ± 2.7 | 22.1 ± 3.2 | 21.3 ± 2.1 | >.9999 | 22.0 ± 1.1 | .6538 |
| Symptoms and signs | | | | | |
| SMPN, n (%) | 18 (100.0) | 9 (100.0) | 9 (100.0) | >.9999 | NA | NA |
| Dysautonomia, n (%) | 17 (94.4) | 8 (88.9) | 9 (100.0) | >.9999 | NA | NA |
| GI symptoms, n (%) | 16 (88.9) | 7 (77.8) | 9 (100.0) | .4706 | NA | NA |
| GU symptoms, n (%) | 12 (66.7) | 5 (55.6) | 7 (77.8) | .6199 | NA | NA |
| Postural dizziness, n (%) | 12 (66.7) | 5 (55.6) | 7 (77.8) | .6199 | NA | NA |
| Skin color changes, n (%) | 8 (44.4) | 4 (44.4) | 4 (44.4) | >.9999 | NA | NA |
| Abnormal glandular secretion, n (%) | 15 (83.3) | 7 (77.8) | 8 (88.9) | >.9999 | NA | NA |
| Chronic cough, n (%) | 8 (44.4) | 5 (55.6) | 3 (33.3) | .6372 | NA | NA |

Note: All the data represent mean ± standard deviation unless otherwise indicated. Abbreviations: BMI, body mass index; GI, gastrointestinal; GU, genitourinary; HC, healthy control; n, numbers; NA, not available; SMPN, sensor-motor polyneuropathy. * and ** represent significant difference at .05 and .01 levels, respectively.
RESULTS

Clinical features of TTR-FAP

The mean age at which these TTR-FAP patients and HCs were examined was 50.4 years (29-64 years) and 47.6 years (26-68 years), respectively. There was no statistically significant difference in age ($p = .5182$) or in the mean height, weight, and body mass index between the two groups (all $p$s > .05) (Table 1).

Of the 18 TTR-FAP patients, 13 initially developed limb paresthesia, followed by other onset symptoms such as diarrhea and/or constipation in 2 patients, sexual dysfunction in 2 patients, and postural dizziness in 1 patient. All patients presented with sensorimotor polyneuropathy, and 17 (94.4%) of them developed autonomic neuropathy, including alternating diarrhea and/or constipation in 16 patients, sexual dysfunction or dysuria in 12 patients, postural dizziness in 12 patients, skin color changes in 8 patients, and abnormal sweating or xerostomia in 15 patients. Specifically, chronic cough was found in 8 patients (44.4%). Nine (50%) patients were early-onset patients (<50 years), and the rest were late-onset ones who were older at the time of examination than early-onset cases and had a lower rate of positive family history. However, there was no statistically significant difference in durations of disease between the early-onset and late-onset patients ($p = .6173$). The two groups were not significantly different in autonomic dysfunction (all $p$s > .05).

Different TTR mutations were identified by gene testing, including Val30Leu mutation in 3 patients, Gly83Arg, Lys35Asn, Val30Met, Glu61Lys, and Ala97Ser mutation in 2 patients, respectively, and Val30Ala, Glu42Gly, Ala36Pro, Phe33Val and Asp38Val mutation in 1 patient, respectively. In clinical staging, 10 of these patients were divided into Coutinho stage I, and the remaining into Coutinho stage II (Table 1).

Ultrasonographic findings

Comparison of VN CSAs between TTR-FAP and HCs

The median (range) CSAs of the bilateral VNs were 3.5 (2.0-6.0) mm$^2$ (right) and 2.5 (1.0-6.0) mm$^2$ (left) in TTR-FAP, and 2.0 (1.0-3.0) mm$^2$ (right) and 1.0 (1.0-2.0) mm$^2$ (left) in HCs, so there was a significant difference between the two groups (all $p$s < .0001) (Figure 1). Furthermore, the median value of mean CSAs of bilateral VNs was higher in Coutinho stage II groups than in stage I groups (3.8 vs. 2.5 mm$^2$, $p = .0074$). However, there was no significant difference between the early-onset and late-onset patients.

![Figure 2](image.jpg)  
**Figure 2** The cross-sectional areas (CSAs) of vagus nerves in transthyretin familial amyloid polyneuropathy (TTR-FAP) patients and controls. n, numbers. ** and *** represent significant difference at 0.01 and 0.001 levels, respectively.
FIGURE 3  Correlation and receiver operator characteristic curve analysis of vagus nerve cross-sectional areas (CSAs) in transthyretin familial amyloid polyneuropathy patients. AUC, area under curve; COMPASS 31, Composite Autonomic Symptom Score 31; NIS, Neuropathy Impairment Score; Norfolk QOL-DN, Norfolk Quality of Life-Diabetic Neuropathy Score. One patient of the longest course of disease was not included in the correlation between CSAs of vagus nerve and the course of disease. Norfolk QOL-DN data for two cases were not available. * and ** represent significant correlation at 0.05 and 0.01 levels, respectively.

The ROC curves of VN CSAs for identifying TTR-FAP from HCs

Based on the results observed in Figure 2, the ROC curve of VN CSAs was used to differentiate between TTR-FAP and HCs. The cutoff value of right and left VN CSAs identified TTR-FAP from HCs were 2.5 and 1.5 mm² and the AUCs were .9395 and .8856, with a high sensitivity (.889 and .889) and specificity (.941 and .765), respectively (Figure 3).

Correlation of mean VN CSAs with clinical measurement scales, cardiac parameters and CSAs of median/ulnar nerves of TTR-FAP

Correlation analysis was conducted between the mean CSAs of bilateral VNs and NIS, Norfolk QOL-DN, COMPASS 31, and duration of disease of TTR-FAP patients. The mean VN CSAs were positively correlated with COMPASS 31 ($r = .5252$, $p = .0252$). It seemed that the larger CSA was accompanied by a longer disease course ($r = .4457$, $p = .0638$) except the longest disease course with a smaller CSA in 1 patient (Table 2). The correlation became stronger without the aforesaid patient ($r = .7203$, $p = .0016$) (Figure 3 and Table 2). Interestingly, mean VN CSAs were positively correlated with IVST ($r = .5426$, $p = .0200$) and LVPWT ($r = .5103$, $p = .0305$). However, mean VN CSAs were not correlated with Serum NT-pro BNP level, NIS, or Norfolk QOL-DN. Nor were mean VN CSAs correlated with either the maximum CSA or mean CSA of median/ulnar nerves (all $p$s > .05) (Table 2).

DISCUSSION

Our early study indicated that autonomic dysfunction was often observed before sensory and motor nerve dysfunction in TTR-FAP. We found that autonomic dysfunction was not significantly different between early- and late-onset disease groups. Other studies indicated that early-onset patients displayed autonomic dysfunction more distinctively than late-onset ones. Most of the patients had a peripheral neuropathy with variable autonomic symptoms. We found that...
The enlargement of periph-
tral nerves of limbs and atrophy of VN were observed in DPN, suggesting that the severity and progression of autonomic neuropathy and sensorimotor neuropathy might not be parallel, indicating that the progression of autonomic neuropathy followed its own pattern that could be better reflected by the CSAs of VN. Niu et al. found a corre-
lation between the CSAs of VN and the maximum/mean CSAs of the median/ulnar nerves in CIDP and CMT1A, which was different from the pattern in TTR-FAP. However, we did not find any correlation in our study, which offered more evidence that the enlargement of VN in TTR-
FAP might have a different pathological mechanism from other periph-
eral neuropathies. Our study found that the CSAs of VN were positively corre-
lated with the course of disease if the patient with the longest course of disease was excluded. It seemed that in the early stage of TTR-FAP, VN increased in size with the prolongation of the course of disease, and then decreased in size in the later stage of the disease. However, because of the limited number of cases, we could not reach a definitive conclusion. What we could do was to recommend VN ultra-
sound as a monitoring index for the progression of TTR-FAP autonomic neuropathy.

Nerve ultrasound, as a noninvasive examination tool, was of great significance for the differential diagnosis of different types of dis-
eases. Patients of late-onset and sporadic TTR-FAP were easily misdi-
agnosed as CIDP, DPN, and other mimics. The enlargement of periph-
eral nerves of limbs and atrophy of VN were observed in DPN, which might be a distinctive trait by which to distinguish TTR-FAP from DPN. Enlargement of VN could be observed in both CIDP and TTR-FAP. However, our previous study explored the difference in nerve ultrasound between CIDP and TTR-FAP in median, ulnar, tibial, sciatic, peroneal, and sural nerves, so we highlighted the dif-
fERENCE in the pattern of nerve enlargement and CSA variability of median nerves between CIDP and TTR-FAP. There was no signi-
ficant difference in the variability of bilateral VN CSAs between TTR-
FAP patients and HCs, suggesting that TTR-FAP patients had relatively symmetrical VN involvement. Future cohort studies could com-
pare the CSAs and the CSA variability of VN between TTR-FAP and CIDP.

This study has several limitations. The sample size of TTR-FAP patients is not big enough, so more subjects will be needed in the future. In addition, the mimics of TTR-FAP and asymptomatic carriers need to be compared with TTR-FAP patients.

In conclusion, VN enlargement is prevalent in TTR-FAP patients. VN ultrasound is a useful tool for evaluation of TTR-FAP.

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