Effects of omalizumab therapy on peripheral nerve functions: short observational study

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Abstract

Introduction: Peripheral neuropathy (PN) is a common neurological condition causing symmetrical and diffuse damage in nerves. The etiology of PN includes systemic diseases, toxic exposure, medications, infections, and hereditary diseases. Omalizumab is a humanized monoclonal anti-IgE antibody that exerts its activity by binding to free IgE in circulation.

Aim: To investigate the relationship between omalizumab and peripheral neuropathy.

Material and methods: The study included 30 patients who underwent omalizumab therapy (Xolair) due to the diagnosis of chronic urticaria. A detailed neurological and physical examination was performed in each patient both before and 3 months after the therapy. Electrophysiological examination was also performed using a Medelec Synergy instrument.

Results: The 30 patients included 8 (26.7%) men and 22 (73.3%) women with a mean age of 37.5 ±14.14 years. No serious side effect of the medication was detected in any patient although local wound irritation occurred in 3 (10%) patients. Moreover, no change occurred in the pre-treatment Neuropathy Symptom Score (NSS) or Neurological Disability Score (NDS) of the patients and no pathological values that could result in neuropathy were observed during motor/sensory nerve conduction. However, significant changes were detected in the sensory and motor components of the nerves with regards to pre- and post-treatment values.

Conclusions: Omalizumab therapy caused no peripheral neuropathy in any of our patients but altered the latency, amplitude, and velocity values of the peripheral nerves.

Key words: chronic urticaria, omalizumab, neuropathy.

Introduction

Peripheral neuropathy (PN) is one of the most common neurological conditions, causing symmetrical and diffuse damage in nerves. The etiology of PN includes systemic diseases, toxic exposure, medications, infections, and hereditary diseases. The most common medications associated with PN include amiodarone, chloroquine, hydralazine, lithium, metronidazole, phenytoin, isoniazid, statins, and vincristine. In addition, biological agents including infliximab and adalimumab have been reported in recent case studies. The prevalence of PN has been reported to be as high as 2.4% in the general population and to be 26.4% in patients with diabetes mellitus [1–4].

Immunoglobulin E (IgE) plays a central role in the pathogenesis of allergic conditions. Therefore, anti-IgE therapies play a key role in the treatment of allergic diseases such as asthma [1–5]. Omalizumab is a humanized monoclonal anti-IgE antibody. Omalizumab exerts its activity by binding to free IgE in circulation, thereby inhibiting the binding of IgE to its high-affinity receptors (FcεRI) found on mast cells and basophils, ultimately reducing the expression of mediators in mast cells. Omalizumab is also an important treatment option particularly for severe asthma and resistant chronic urticaria [6–8].

On the other hand, although histamine is the most important mediator expressed in mast cells, neuropeptides such as nerve growth factor (NGF) are also expressed in these cells [9]. A previous study reported that...
mast cell activation led to an increase in the production and secretion of neuropeptides and the excitability of sensory nerves [10]. A recent study evaluated the effectiveness of omalizumab at 52 weeks and showed that omalizumab is a safe drug although it had several side effects including headache, injection site reaction, myalgia, lethargy, nausea, dizziness, weight gain, and arthralgia [11]. However, another study showed that omalizumab therapy resulted in optic neuritis when used for the treatment of bronchial asthma in 2 patients with Churg-Strauss syndrome [12]. Similarly, Lieberman et al. compared omalizumab therapy with placebo therapy and reported that local skin reactions occurred in 44% of the patients treated with omalizumab [13].

In our patients, we also performed omalizumab therapy for the treatment of chronic urticaria. However, the complaints of pain and weakness in the extremities gradually increased in our patients and thus we could not be sure whether these complaints resulted from local irritation or a neurological condition caused by omalizumab, mainly because the drug was administered in two separate infusions with 150 mg flacons. Moreover, these conditions may be a result of peripheral nerve injury caused by the inhibition of mast cells that leads a reduction in the expression of neuromediators.

Aim

In this study, we aimed to evaluate the relationship between omalizumab and peripheral neuropathy.

Material and methods

The study included 30 patients who underwent omalizumab therapy (Xolair) due to the diagnosis of chronic urticaria. Age, gender, socioeconomic status, and family history were recorded for each patient. Omalizumab was subcutaneously administered at 300 mg/day for 28 days (total 4 times: 1, 29, 57, 85 days) in the Dermatology Department. To determine the presence of other factors that may affect peripheral nerve function, additional tests were performed, including complete blood count, sedimentation rate, liver and kidney function tests, urine analysis, thyroid hormones, vitamin B12 level, folic acid level, and serologic tests. In addition, neuro-radiological imaging was performed as needed. Exclusion criteria included neurological symptoms and signs, diabetes mellitus, connective tissue disease, hepatic, renal, and thyroid diseases, amyloidosis, heart failure, alcohol abuse, corticosteroid use, cervical disc hernia, and malignancy. A detailed neurological and physical examination was performed in each patient before and three months after the therapy. Neurological symptoms were scored using the Neuropathy Symptom Score (NSS) and Neurological Disability Score (NDS). Following the neurological examination, electrophysiological examination was performed using a Medelec Synergy instrument (Oxford Instruments, Surrey, UK) with standard neurographic procedures, and the results were evaluated according to the American Diabetes Association (ADA) Diabetic Neuropathy Guidelines in the Neurology Department [14]. The measurements were performed 24 h before and 90 days after omalizumab therapy. Room temperature was kept at 22–24°C and the temperature of the extremity was kept at 34°C and it was heated as needed. Nerve conduction tests were performed in two motor and two sensory nerves (median and ulnar nerves) in the upper extremities and in two motor (tibial and common peroneal nerve) and two sensory nerves (sural and peroneal sensory nerves) in the lower extremities. Pre- and post-treatment latency (ms), amplitude (mV), and velocity (m/s) values were compared for each nerve. Nerve conduction velocity was measured using the orthodromic method and nerve conduction was performed at supramaximal intensity to achieve the highest amplitude. Presence of an axonal pathology and demyelination in the nerves was defined as decreased sensory/motor nerve action potential amplitude and slowing of sensory/motor nerve conduction velocity. Polyneuropathy was defined as the presence of two or more abnormalities detected in electrophysiological examination. The study was approved by the local ethics committee and informed consent was obtained from each patient (Number: YYU: 2017/02).

Statistical analysis

Data were analyzed using IBM SPSS for Windows, Version 20.0 (Armonk, NY: IBM Corp.). Normal distributions of variables were determined by histogram and/or by the Kolmogorov-Smirnov/Shapiro-Wilk test. Descriptive statistics were expressed as mean, standard deviation (SD), median, minimum, and maximum. Numerical variables were compared using the paired sample t-test for data with normal distribution and the Wilcoxon signed-rank test for data with non-normal distribution. Spearman’s correlation coefficient was used to assess the correlation between variables. A p-value of < 0.05 was considered significant.

Results

The 30 patients included 8 (26.7%) men and 22 (73.3%) women with a mean age of 37.5 ±14.14 years (Table 1). No serious side effect of the medication was observed in any patient although local wound irritation occurred in 3 (10%) patients. No change occurred in the pre-treatment NSS and NDS scores of the patients. Moreover, no pathological values that could result in neuropathy were observed during motor/sensory nerve conduction. Nevertheless, significant changes were detected in the sensory and motor components of the nerves with regards to pre- and post-treatment values.
A comparison of pre- and post-treatment latency, amplitude, and velocity values of the sensory component of the median nerves indicated no significant difference between the pre- and post-treatment latency and velocity values ($p > 0.05$). However, post-treatment amplitude values (mean: $23.03 \pm 4.71$) were significantly lower than pre-treatment values (mean: $24.61 \pm 5.55$) ($p = 0.024$) (Table 2).

No significant difference was found between the pre- and post-treatment latency and velocity values of the sensory component of the ulnar nerves ($p > 0.05$). Nevertheless, post-treatment amplitude values (mean: $18.71 \pm 5.34$) were significantly lower than pre-treatment values (mean: $20.11 \pm 5.61$) ($p = 0.030$) (Table 3).

In the sensory component of the sural nerves, no significant difference was found between the pre- and post-treatment amplitude values ($p > 0.05$). However, post-treatment latency values (mean: $2.12 \pm 0.24$) were significantly higher than pre-treatment values (mean: $1.96 \pm 0.21$) ($p = 0.009$), whereas post-treatment velocity values (mean: $46.34 \pm 3.60$) were significantly lower than pre-treatment values (mean: $48.14 \pm 7.09$) ($p = 0.001$) (Table 4).

In the sensory component of the superficial peroneal nerves, no significant difference was found between pre- and post-treatment latency values ($p > 0.05$), whereas post-treatment amplitude and velocity values (mean: $14.00 \pm 3.02$ and $47.20 \pm 3.36$, respectively) were significantly lower than pre-treatment values (mean: $16.45 \pm 4.05$ and $49.34 \pm 3.36$, respectively) ($p < 0.001$ for both) (Table 5).

On the other hand, a comparison of pre- and post-treatment latency, amplitude, and velocity values of the motor component of the median nerves indicated no significant difference between the pre- and post-treatment latency values ($p > 0.05$) although post-treatment amplitude and velocity values (mean: $8.40 \pm 1.54$ and $57.97 \pm 4.42$, respectively) were significantly lower than pre-treatment values (mean: $9.21 \pm 2.08$ and $59.42 \pm 4.61$, respectively) ($p = 0.007$ and $0.049$, respectively) (Table 6).

Moreover, no significant difference was found between the pre- and post-treatment latency and amplitude values of the motor component of the ulnar nerves ($p > 0.05$). However, post-treatment velocity values (mean: $57.62 \pm 4.13$) were significantly lower than pre-treatment values (mean: $59.04 \pm 4.02$) ($p = 0.002$).

In the motor component of the superficial peroneal nerves, post-treatment latency values (mean: $3.90 \pm 0.45$) were significantly higher than pre-treatment values.

### Table 1. Characteristics of patients

| Parameter            | N   | %   |
|----------------------|-----|-----|
| Gender:              |     |     |
| Male                 | 8   | 26.67 |
| Female               | 22  | 73.33 |
| Marital status:      |     |     |
| Married              | 23  | 76.67 |
| Single/other         | 7   | 23.33 |
| Education status:    |     |     |
| Primary school       | 6   | 20.00 |
| Middle school        | 8   | 26.67 |
| High school          | 9   | 30.00 |
| University           | 7   | 23.33 |
| Income status:       |     |     |
| Low income           | 15  | 50.00 |
| Middle income        | 11  | 36.67 |
| High income          | 4   | 13.33 |
| Age*                 | 37.50 ±14.14 | 35.50 |
| Disease duration [months]* | 12.67 ±8.05 | 10.50 |
| Cumulative dose [for each patient] | 1200 mg |

*Results are expressed as mean ± SD and median data instead of N and %.

### Table 2. Comparison of pre- and post-treatment latency, amplitude, and velocity values of the sensory component of the median nerves

| Variable         | Mean ± SD | Median | Minimum | Maximum | P-value |
|------------------|-----------|--------|---------|---------|---------|
| Latency-pre      | 2.17 ±0.23| 2.13   | 1.75    | 2.75    | 0.278*  |
| Latency-post     | 2.23 ±0.31| 2.13   | 1.80    | 3.00    |         |
| Amplitude-pre    | 24.61 ±5.55| 25.50  | 13.50   | 35.00   | 0.024*  |
| Amplitude-post   | 23.03 ±4.71| 23.10  | 13.50   | 31.90   |         |
| Velocity-pre     | 55.67 ±5.81| 55.60  | 40.00   | 63.40   | 0.821*  |
| Velocity-post    | 55.28 ±5.24| 55.95  | 42.00   | 63.90   |         |

*Paired samples t-test, *Wilcoxon signed-rank test, pre – pre-treatment, post – post-treatment.
Table 3. Comparison of pre- and post-treatment latency, amplitude, and velocity values of the sensory component of the ulnar nerves

| Variable     | Mean ± SD  | Median | Minimum | Maximum | P-value |
|--------------|------------|--------|---------|---------|---------|
| Latency-pre  | 1.89 ±0.19 | 1.90   | 1.50    | 2.45    | 0.436*  |
| Latency-post | 1.91 ±0.22 | 1.83   | 1.60    | 2.40    |         |
| Amplitude-pre| 20.11 ±5.61| 18.95  | 11.90   | 34.00   | 0.030*  |
| Amplitude-post| 18.71 ±5.34| 17.45  | 12.00   | 31.00   |         |
| Velocity-pre | 54.76 ±2.99| 54.90  | 50.00   | 61.10   | 0.209*  |
| Velocity-post| 53.94 ±3.60| 52.75  | 44.90   | 60.60   |         |

*aPaired samples t-test, bWilcoxon signed-rank test, pre – pre-treatment, post – post-treatment.

Table 4. Comparison of pre- and post-treatment latency, amplitude, and velocity values of the sensory component of the sural nerves

| Variable     | Mean ± SD  | Median | Minimum | Maximum | P-value |
|--------------|------------|--------|---------|---------|---------|
| Latency-pre  | 1.96 ±0.21 | 2.00   | 1.55    | 2.42    | 0.009*  |
| Latency-post | 2.12 ±0.24 | 2.10   | 1.55    | 2.55    |         |
| Amplitude-pre| 15.39 ±3.30| 14.55  | 9.40    | 22.20   | 0.053*  |
| Amplitude-post| 14.48 ±3.32| 14.20  | 8.00    | 21.80   |         |
| Velocity-pre | 48.14 ±7.09| 48.85  | 16.80   | 58.10   | 0.001*  |
| Velocity-post| 46.34 ±3.60| 45.80  | 41.90   | 56.90   |         |

*aPaired samples t-test, bWilcoxon signed-rank test, pre – pre-treatment, post – post-treatment.

Table 5. Comparison of pre- and post-treatment latency, amplitude, and velocity values of the sensory component of the superficial peroneal nerves

| Variable     | Mean ± SD  | Median | Minimum | Maximum | P-value |
|--------------|------------|--------|---------|---------|---------|
| Latency-pre  | 2.00 ±0.28 | 2.00   | 1.50    | 2.55    | 0.497   |
| Latency-post | 2.05 ±0.28 | 2.05   | 1.60    | 2.90    |         |
| Amplitude-pre| 16.45 ±4.05| 16.20  | 9.20    | 23.50   | < 0.001 |
| Amplitude-post| 14.00 ±3.02| 14.30  | 9.00    | 21.80   |         |
| Velocity-pre | 49.34 ±4.61| 48.25  | 41.90   | 55.90   | 0.002   |
| Velocity-post| 47.20 ±3.36| 47.45  | 41.30   | 55.90   |         |

*aPaired samples t-test, bWilcoxon signed-rank test, pre – pre-treatment, post – post-treatment.

Table 6. Comparison of pre- and post-treatment latency, amplitude, and velocity values of the motor component of the median nerves

| Variable     | Mean ± SD  | Median | Minimum | Maximum | P-value |
|--------------|------------|--------|---------|---------|---------|
| Latency-pre  | 2.76 ±0.35 | 2.73   | 2.15    | 3.70    | 0.052*  |
| Latency-post | 2.91 ±0.47 | 2.80   | 2.20    | 3.95    |         |
| Amplitude-pre| 9.21 ±2.08 | 8.70   | 6.60    | 15.80   | 0.007*  |
| Amplitude-post| 8.40 ±1.54 | 8.10   | 5.70    | 13.50   |         |
| Velocity-pre | 59.42 ±4.61| 60.15  | 51.00   | 69.40   | 0.049*  |
| Velocity-post| 57.97 ±4.42| 58.60  | 50.00   | 64.90   |         |

*aPaired samples t-test, bWilcoxon signed-rank test, pre – pre-treatment, post – post-treatment.
Advances in Dermatology and Allergology

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Alvarez-Lario syndrome with omalizumab therapy [21]. On the other hand,
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zumab on the nerves. Jachiet number of studies investigating the effect of omali-
umb in patients with chronic urticaria. Kim reviewed more
peripheral neuropathy. However, further studies are needed to shed light on our findings.

Discussion

The results indicated that omalizumab did not cause peripheral neuropathy but altered the latency, amplitude,
to trigger the auto-immune process, as observed in a patient described by Kalteren et al. In addition, depending on the finding that
omalizumab changed the latency, amplitude, and velocity values of peripheral nerves in our patients, we be-
ial, with a growing side-effect profile. In this study, we
evaluated the relationship between omalizumab and peripheral neuropathy.

We evaluated the effect of omalizumab in patients with chronic urticaria. Kim et al. also evaluated the ef-
effect of omalizumab in patients with chronic spontaneous urticaria and reported that 61.75% of the patients were
women. Similarly, women also constituted the majority of our patients (73.3%), which implies that chronic urti-
caria has a female preponderance [15].

Omalizumab has been shown to be a safe drug in numerous studies. However, a number of side effects have been associated with omalizumab, including anaphylaxis, urticaria, eosinophilic granulomatosis with polyangiitis, susceptibility to parasitic infections, injection site reactions, cardiovascular diseases, and serum sickness [13, 16–20]. On the other hand, a previous study reported that headache and disturbance of sleep were the most common neurological side effects of omalizumab [19]. Similarly, Corren et al. reviewed more than 7,500 patients undergoing omalizumab therapy and found that headache was the most common neu-
rolological side effect and also noted that omalizumab led to musculoskeletal disturbances including low back pain, arthralgia, pain in the extremities, and myalgia [17]. In our study, no complaint of headache was found in any patient, which could be ascribed to the small pa-
tient series in our study.

Literature reviews indicate that there are a limited number of studies investigating the effect of omali-
umb on the nerves. Jachiet et al. reported that omali-
zumab therapy resulted in optic neuritis in 2 patients [12]. In contrast, Kalteren et al. evaluated a patient with ocular myasthenic syndrome and reported that all the symptoms were resolved after the treatment of the syn-
drome with omalizumab therapy [21]. On the other hand, Alvarez-Lario et al. evaluated the effectiveness of biologi-
cal treatment and reported that the side effects of the
treatment resulted in peripheral neuropathy associated with Guillain-Barré syndrome in 21 patients. The authors
considered that peripheral neuropathy resulted from the increased susceptibility to infections caused by biological agents [22]. As shown in these studies, omalizumab typi-
cally increases susceptibility to infections. However, no infection associated with omalizumab was observed in our patients. On the other hand, although no peripheral neuropathy occurred in any of our patients, we consider
that omalizumab has the potential to affect peripheral nerves since it has been shown to increase susceptibility to infections, to alter the growth factors and neuropeptides expressed in mast cells, and to trigger the auto-
immune process, as observed in a patient described by Kalteren et al. In addition, depending on the finding that
omalizumab changed the latency, amplitude, and velocity values of peripheral nerves in our patients, we be-

Our study was limited since it was a single-center study and had a relatively small number of patients. Moreover, since we assessed the neurological symptoms of the patients before and three months after the treat-
ment, different outcomes could have been detected if the symptoms had also been assessed at 1 year after the treatment.

Conclusions

Omalizumab therapy is becoming gradually popu-
lar, with a growing side-effect profile. In this study, we investigated the relationship between omalizumab and neuropathy. However, further studies are needed to shed light on our findings.

Conflict of interest

The authors declare no conflict of interest.

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