Parsonage‐Turner syndrome following COVID-19 vaccination

INTRODUCTION
Parsonage‐Turner syndrome (PTS), also known as neuralgic amyotrophy, is an acute idiopathic brachial neuritis, typically characterised by acute onset of excruciating pain followed by weakness and wasting in the upper limb.¹ Antecedent events such as infection, exercise, trauma, surgery and vaccination are reported in approximately 50% of affected individuals.¹ PTS has been reported following COVID-19 vaccination, but the current literature is limited to several case reports and a passive reporting system.² ³ Herein, we report on the clinical, radiological and laboratory features of 12 cases with PTS post COVID-19 vaccination.

METHODS
We reviewed medical records of the patients who were diagnosed with PTS following COVID-19 vaccination in three referral hospitals (Seoul, South Korea) between June and October 2021. We collected and analysed the detailed clinical information as follows: the type and order (in case of different types) of COVID-19 vaccine, laterality of symptom presentation, timeline regarding the vaccination, symptom onset and nadir, clinical presentation, motor grade at nadir, the results of electrodiagnosis, brachial plexus MRI, cerebrospinal fluid (CSF) analysis, treatment regimen and the outcomes.

RESULTS
We identified 12 patients (7 men and 5 women) who developed PTS after a receipt of COVID-19 vaccine. Clinical features of the patients are summarised in table 1 (and online supplemental figure). Age ranged between 23 and 81 (average 51). Vaccination was the only possible trigger in all cases. Six patients had received adenoviral vector-based vaccines (four received AstraZeneca and two received Janssen), and the others had mRNA-based vaccines (five had Pfizer and one had Moderna).

All but two developed PTS after receipt of the first dose of COVID-19 vaccine. The interval from the vaccination to symptom onset and nadir ranged between 2 days and 16 days (median 6.5) and between 5 days and 55 days (median 20.5), respectively. Disease severity at nadir varied across patients, with Medical Research Council grade of the weakest muscles ranging from 2 to 5. Electrodiagnostic studies revealed abnormalities consistent with brachial neuritis in most patients.

Intriguingly, PTS occurred at the same side of vaccine injection in all but two cases (contralateral in patient 8 and bilateral in patient 12). Notably, MRI or sonographic evaluations revealed prominent ipsilateral axillary and/or cervical lymph nodes in seven of eight patients (87.5%) (online supplemental figure). CSF analysis showed albuminocytological dissociation in all three tested patients (patients 1, 3 and 12). We administered oral or intravenous corticosteroid in all patients but three: two patients showed rapid clinical improvement (patients 5 and 7), and one patient refused to receive the treatment.

| No | Sex/age (years) | Vaccine (dose) | Laterality | Days from vaccination to symptom onset/nadir | Weakest muscle strength at nadir | MRI | CSF analysis* | Treatment | Outcome |
|----|----------------|----------------|------------|---------------------------------|---------------------------------|-----|---------------|-----------|---------|
| 1  | M/31           | Janssen        | Ipsilateral| 6/7                             | Prominent ipsilateral axillary and cervical lymph nodes | WBC 0, protein 70 | None | Full recovery by week 1 |
| 2  | M/37           | Janssen        | Ipsilateral| 14/14                           | Normal                          | ND  | Oral prednisolone, gabapentin | Near-full recovery by week 10 |
| 3  | M/71           | AstraZeneca (first dose) | Ipsilateral| 16/35                           | Signal changes and enlargement from the C8 root to the inferior trunk, prominent ipsilateral cervical and axillary lymph nodes | WBC 2, protein 57 | Oral prednisolone, gabapentin | Poor recovery by week 15 |
| 4  | M/63           | AstraZeneca (first dose) | Ipsilateral| 14/14                           | ND                              | ND  | None | Poor recovery by week 4, lost to follow-up thereafter |
| 5  | F/65           | AstraZeneca (first dose) | Ipsilateral| 5/6                             | Prominent ipsilateral axillary lymph nodes | ND  | None | Full recovery within 2 months |
| 6  | M/61           | AstraZeneca (second dose) | Ipsilateral| 2/3                             | Signal changes and enlargement from the C8 root to the inferior trunk and medial cord, prominent cervical and axillary lymph nodes | ND  | Oral prednisolone | Partial recovery by month 5 |
| 7  | F/31           | Cross-vaccination (AstraZeneca and then Pfizer)² | Ipsilateral| 2/10                            | Prominent ipsilateral axillary and cervical lymph nodes | ND  | None | Full recovery by week 3 |
| 8  | F/50           | Pfizer (first dose) | Contralateral| 4/16                           | Normal                          | ND  | NSAIDs, fentanyl patch, IVMP | Good response to IVMP |
| 9  | M/58           | Pfizer (first dose) | Ipsilateral| 5/30                            | ND                              | ND  | Oral prednisolone, pregabalin | Poor recovery by week 8 |
| 10 | F/23           | Pfizer (first dose) | Ipsilateral| 10/11                           | ND (ipsilateral axillary lymphadenopathy in ultrasonography) | ND  | Oral prednisolone, pregabalin | Partial recovery within week 6 |
| 11 | F/81           | Pfizer (first dose) | Bilateral  | 15 NA                           | IV                              | ND  | Pregabalin, nortriptyline, NSAIDs | Poor recovery by month 6 |
| 12 | M/39           | Moderna (first dose) | Ipsilateral| 7/14                            | Prominent ipsilateral axillary lymph nodes | WBC 1, protein 76 | IVMP followed by oral prednisolone, gabapentin | Poor recovery by week 8 |

*Values were expressed as cells/μL (WBC) and mg/dl (protein).
²Parsonage-Turner syndrome occurred 2 days after Pfizer vaccination following initial AstraZeneca vaccination.
³CSF, cerebrospinal fluid; F, female; IVMP, intravenous methylprednisolone; M, male; MRC, Medical Research Council; ND, not done; NSAIDs, non-steroidal anti-inflammatory drugs; WBC, white blood cell.
DISCUSSION

Vaccination is one of the known potential triggers of PTS, with case series and reports for tetanus, smallpox, human papillomavirus, influenza and, recently, COVID-19 vaccines. To date, there have been seven publications reporting 10 cases of PTS following COVID-19 vaccination (online supplemental table). While most of the reported cases (8/10) received mRNA vaccines, our observations implicate the adenoviral vector-based vaccine as well in 6 of 12 cases. The time interval between vaccination and PTS onset ranged from 2 to 16 days, which is in line with previous reports and coincided with the expected time course of the immune response against COVID-19 vaccines. Similar to previous reports, we found that PTS post-COVID-19 vaccination mostly takes the classic form, but also observed atypical phenotypes of pure sensory or painless motor-predominant form in two cases.

Intriguingly, we found that PTS occurred on the same side of vaccination in all except two cases. Combined with all previously reported cases, the rate of ipsilateral PTS reaches 77.3% (17 of 22 cases). Meanwhile, we noted an unexpectedly high frequency of ipsilateral reactive lymphadenopathy in our cases (87.5%). Although it has not been addressed in the context of PTS, reactive lymphadenopathy is reportedly a frequent finding at imaging with an incidence of up to 53% with the COVID-19 mRNA vaccines in patients with breast cancer. Taken together, we propose the need for further research to investigate whether these local immune reactions are involved in the pathogenesis of ipsilateral brachial neuritis or whether they are simply observed together by chance.

It should be emphasised that we cannot establish a robust causal relationship between PTS and COVID-19 vaccination with case series. It is also worth mentioning that the benefits of COVID-19 vaccination far outweigh the potential risks. Corroborating the findings of previous case reports, our observations suggest that COVID-19 vaccines may be associated with PTS characterised by ipsilateral occurrence and possibly accompanied by reactive lymphadenopathy. Further studies are warranted to assess the causality and significance of the ipsilateral association.

Young Gi Min 1,2, Jee-Eun Kim 3, Ji Young Hwang 4, Je-Young Shin 1, Jung-Joon Sung 1, Yoong-Ho Hong 5
1 Neurology, Seoul National University Hospital, Seoul, South Korea
2 Translational Medicine, Seoul National University College of Medicine, Seoul, South Korea
3 Neurology, College of Medicine, Ewha Womans University, Seoul, South Korea
4 Radiology, College of Medicine, Ewha Womans University, Seoul, South Korea
5 Neurology, Neuroscience Research Institute, Medical Research Council, Seoul National University College of Medicine, Seoul Metropolitan Government Boramae Medical Center, Seoul, South Korea

Correspondence to Professor Yoong-Ho Hong, Neurology, Neuroscience Research Institute, Medical Research Council, Seoul National University College of Medicine, Seoul Metropolitan Government Boramae Medical Center, Seoul 07061, South Korea; nhong@ehwa.ac.kr

Twitter Yoong-Ho Hong @YoongHo

Contributors YGM, J-EK and Y-HH conceptualised, designed the study and wrote the manuscript. YGM, JYH and J-EK collected data. YGM and J-EK prepared the figure. J-YS and J-JS interpreted the data. Y-HH supervised the work.

Funding This work was supported by a focused clinical research grant-in-aid (03-2016-0300) from the Seoul National University Hospital Research Fund.

Competing interests None declared.

Patient consent for publication Consent obtained directly from patient(s).

Ethics approval This study involves human participants and was approved by the institutional review boards (IRBs) of Seoul National University Hospital (IRB 2110-049-1261), Seoul Metropolitan Government Boramae Medical Center (IRB 30-2021-122) and Ewha Womans University Seoul Hospital (IRB 2021-09-021). The participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Supplemental material This content has been supplied by the author(s). It has not been vetted by the figure supplier. Directly from patient(s).

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non-Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

© Author(s) (or their employer(s)) 2022. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

Additional supplemental material is published online only. To view, please visit the journal online (http://dx.doi.org/10.1136/jnnp-2021-328182).

YGM and J-EK contributed equally.

YGM and J-EK are joint first authors.

To cite Min YG, Kim J-E, Hwang JY, et al. J Neurol Neurosurg Psychiatry 2022;93:1231–1232.

Received 13 October 2021
Accepted 23 February 2022
Published Online First 6 April 2022
J Neurol Neurosurg Psychiatry 2022;93:1231–1232.

doi:10.1136/jnnp-2021-328182

ORCID iDs
Young Gi Min http://orcid.org/0000-0002-8091-7585
Jee-Eun Kim http://orcid.org/0000-0002-3811-3479
Yoong-Ho Hong http://orcid.org/0000-0002-3325-6358

REFERENCES
1 van Allen N, van Engelen BGM, Allen N, et al. The clinical spectrum of neuralgic amyotrophy in 246 cases. Brain 2006;129:438–50.
2 Queller SC, Towbin AJ, Milani C, et al. Parsonage-Turner syndrome following COVID-19 vaccination: Mr neurography. Radiology 2022;302:211374.
3 Noseda R, Ripellino P, Ghidossi S, et al. Reporting of acute inflammatory neuropathies with COVID-19 vaccines: subgroups Disproportionality analyses in VigiBase. Vaccines 2021;9:1022.
4 Cacciavillani M, Salvaglocio A, Bianci C. Pure sensory neuralgic amyotrophy in COVID-19 infection. Muscle Nerve 2021;63:E7–E8.
5 Garreffa E, Hamad A, O’Sullivan CC, et al. Regional lymphadenopathy following COVID-19 vaccination: literature review and considerations for patient management in breast cancer care. Eur J Cancer 2021;159:38–51.