Introduction
Accessory deep peroneal nerve (ADPN) is the most common anomalous innervation present in the lower limb.

Aim
The aim of this study was to determine the prevalence of ADPN electrophysiologically in a sample of healthy Egyptian individuals.

Subjects and methods
This cross-sectional study included 200 lower limbs from 100 [56 (56%) women and 44 (44%) men] Egyptian apparently healthy volunteers. Motor nerve conduction studies for the peroneal nerve and ADPN were done.

Results
ADPN was found in 20 (10%) lower limbs of 17 (17%) subjects. There was no statistically significant difference between the occurrence of ADPN in women versus men ($P=0.797$). ADPN was found in 11 (55%) right lower limbs and in nine (45%) left lower limbs. There was no statistically significant difference between the occurrence of ADPN in right lower limbs versus left lower limbs among subjects with ADPN ($P=0.637$). It was present bilaterally in three (17.65%) subjects. There was no statistically significant difference between the occurrence of bilateral ADPN in women versus men ($P=0.761$). Among the 14 (82.35%) subjects with unilateral ADPN, it was present in the right side in eight (57.14%) subjects. There was no statistically significant difference between the occurrence of unilateral ADPN in the right side versus left side ($P=0.579$).

Conclusion
This study demonstrated that ADPN prevalence in the referred Egyptian sample through electrodiagnostic studies of lower limbs was 17%, with no sex nor side difference. Recognition of ADPN is essential for proper interpretation of lower limbs electrophysiological data.

Keywords:
accessory deep peroneal nerve, anomalous innervations, Egyptian population

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The research was explained thoroughly, then an informed consent was given by each participant. The Ethics Committee of the Faculty of Medicine, Alexandria University, Egypt had approved the study.

Demographic data were collected from all studied subjects. Neurological examination was done for all of them.

Electrophysiological studies were conducted on a Nihon Kohden Neuropack S1 MEB-9400 unit with a two-channel evoked potential/EMG measuring system (Nihon Kohden Corporation, Tokyo, Japan). Temperature of the skin (at the site of the recording electrodes) was maintained at around 32–34°C by means of an infrared lamp. The ground electrode was placed between the stimulation site proximally and the recording electrodes distally [1]. A measurement tape was used for measuring the conduction distances with a precision of 1 mm.

The following parameters were applied for the motor nerve conduction studies: The filter bandwidth was 10 Hz–10 kHz. The pulse duration was 0.2 ms. The current production ability of the bipolar stimulator was 50 mA. The compound muscle action potentials (CMAP) amplitude was measured from the baseline to the negative peak expressed in millivolts. Supramaximal stimulation was ensured. The electrophysiological studies were done by caution. It was essential to make sure that the change in the CMAP amplitude obtained between distal and proximal sites of stimulation was not due to technical factors such as submaximal stimulation and co-stimulation of the nearby nerves [1].

The peroneal motor nerve conduction study and ADPN motor nerve conduction study recording the EDB muscle were done as the following: the active recording surface disc electrode was placed over the EDB muscle belly on the dorsolateral aspect of the foot and the reference surface disc electrode was placed over the dorsal aspect of the metatarsophalangeal joint of the fifth toe (Figs 2 and 3) [1].

Electrical stimulation of the peroneal nerve was done at the following sites: (i) distal stimulation: at 9 cm proximal to the active recording electrode at the ankle slightly lateral to the tibialis anterior tendon; (ii) proximal stimulation at two sites: the first site is below fibular head on the lateral calf at about two-finger breadths inferior to the fibular head, and the second site is at the lateral popliteal fossa adjacent to the external hamstring tendons at a distance of 10 cm proximal to the below fibular head site of stimulation (Fig. 2). The sweep speed was 5 ms/division and the sensitivity was 5 mV/division. The CMAP amplitude was taken for analysis [1].

Criteria for suspecting the presence of ADPN electrophysiologically included the following [1,11]:

1. The peroneal CMAP amplitude recorded from the EDB muscle was higher at the below fibular head and lateral popliteal fossa sites of stimulation than that obtained at the ankle site of stimulation.
(2) Absent peroneal CMAP while recording from the EDB muscle, stimulating the ankle site, with elicited peroneal CMAP from EDB muscle stimulating proximally.

Electrical stimulation of the ADPN was done behind the lateral malleolus for all the participated subjects (Fig. 3). The sweep speed was 5 ms/division and the sensitivity was 1–2 mV/division. The CMAP amplitude was taken for the analysis [1,12,13]. An elicited response is a confirmation for the presence of ADPN [13].

The ADPN was classified electrophysiologically as the following [11,13]:

1. ADPN with partial innervation of the EDB muscle: The ADPN supplies the EDB muscle partially. The peroneal CMAP amplitude recorded at the EDB muscle at the ankle stimulation site is lower than the CMAP amplitude obtained at proximal sites of stimulation. This is associated with ADPN CMAP recorded from the EDB muscle.

2. ADPN with total (exclusive) innervation of the EDB muscle: The ADPN supplies the EDB muscle totally. There is no peroneal CMAP elicited during recording of the EDB muscle at the ankle stimulation site, in spite of the presence of CMAP obtained at proximal sites of stimulation. On the other hand, the ADPN CMAP while stimulating behind the lateral malleolus is recorded from the EDB muscle.

Statistical analysis of the results were done by using the statistical package of the social sciences (SPSS version 17) software (University of Cambridge computing service, London, United Kingdom) [14]. Descriptive measures including count, frequency, minimum, maximum, mean, and standard deviation (SD) were used. Analytic measures included Pearson’s \( \chi^2 \)-test and Fisher’s exact when required. Statistical significance was assigned to any \( P \) value less than 0.05.

Results
The study included 200 lower limbs that were obtained from 100 [56 (56%) women and 44 (44%) men] Egyptian apparently healthy volunteers. Their mean age was 39.19 ± 13.84 years (ranged: 18–75 years). Electrophysiologically, ADPN was found in 20 (10%) lower limbs of 17 (17%) subjects. Among them, it was present in ten (58.82%) women. There was no statistically significant difference between the occurrence of ADPN in women versus men (\( \chi^2 = 0.066, P = 0.797 \)). ADPN occurred in 11 (55%) right lower limbs and in nine (45%) left lower limbs. There was no statistically significant difference between the occurrence of ADPN in right lower limbs versus left lower limbs among subjects with ADPN (\( \chi^2 = 0.222, P = 0.637 \)).

Among subjects with ADPN, it was present bilaterally in three (17.65%) subjects which represent 3% of the total
number of subjects included in this study. It was present in two (66.67%) women. There was no statistically significant difference between the occurrence of bilateral ADPN in women versus men ($\chi^2=0.093$, $P=0.761$).

Among the 14 subjects (82.35%) with unilateral ADPN, it was present in the right side in eight (57.14%) subjects. There was no statistically significant difference between right versus left sides among subjects with unilateral ADPN ($\chi^2=0.307$, $P=0.579$).

All the 17 (100%) subjects with positive ADPN in our study presented with ADPN with partial innervation of the EDB muscle (Fig. 4). No one showed the ADPN with total innervation of the EDB muscle.

Discussion

The ADPN arises from the superficial peroneal nerve as a continuation of the muscular branch supplying the peroneus longus and peroneus brevis muscles (Fig. 1). It descends in the lateral compartment of the leg along the posterior border of the peroneus brevis muscle. At the ankle region, it passes behind the lateral malleolus in close relation to the sural nerve and deep into the peroneus brevis tendon to reach the dorsum of the foot [10,15]. In this region, it gives deep sensory supply to the ankle joint and the surrounding ligaments and tendons. Also, it gives motor branch to supply the lateral aspect of EDB muscle [15]. However, total innervation of the EDB muscle by the ADPN could occur [11,15–17].

Peripheral nerves anomalous innervations are important aspects in routine neurophysiological assessment of any patient. Unrecognition of these anomalous innervations can be mistaken for technical pitfalls or for actual pathology [1,18]. It is important to take into consideration the anatomic variations in the innervations of muscles during electrophysiological assessment of nerves [4,19].

The prevalence of ADPN in a sample of Egyptian subjects was 17% electrophysiologically (10% of the examined lower limbs). The prevalence of ADPN in this study was within the range of prevalence of ADPN present in other studies which varies from 12 to 35% among the studied subjects (8.2–27% of the studied lower limbs) [10]. It was found that there is a wide variation of prevalence of ADPN among different studies [2,12,13,20,21]. A meta-analysis study assessed the prevalence of ADPN and found the following: (i) the overall pooled prevalence was 18.8% of lower limbs; (ii) the electrophysiological pooled prevalence was 13.6%; and (iii) the anatomical pooled prevalence was 39.3% [2]. This could be explained by the differences between studies regarding the studied population and the techniques used in the assessment of ADPN, whether anatomical or electrophysiological studies [2]. The anatomical studies showed a higher prevalence of ADPN than that obtained by electrophysiological studies. It was found that when the ADPN supplies the EDB muscle by a motor branch is the only situation that can be detected electrophysiologically. However, if the ADPN has no motor supply to the EDB muscle, it could not be detected except anatomically [2,10]. It was reported that the pooled prevalence of ADPN with motor supply to the EDB muscle was 16.3% anatomically, which is approximately equal to the electrophysiological pooled prevalence of ADPN [2].

In this research, there was no statistically significant difference between women and men as regards the
frequency of ADPN. This was in agreement with other studies [12,13]. This could be due to the autosomal dominant inheritance of ADPN. It was reported to be about three times more common in the family members of persons who had ADPN [8,20].

In this study, there was no statistically significant difference between the occurrence of ADPN in right lower limbs versus left lower limbs among subjects with ADPN. This was in accordance with other studies [2,10,12,13,20]. In this study, unilateral ADPN was present in 82.35% of the subjects with ADPN. This was in agreement with previous studies which reported that unilateral ADPN was more common than bilateral ADPN [2,12,13].

Regarding the electrophysiological forms of ADPN, ADPN with partial innervation of the EDB muscle was the only form to be present. It is the form in which the ADPN supplies the lateral part of the EDB muscle, while its medial part is supplied by the deep peroneal nerve. All our 17 subjects with recorded ADPN had ADPN with partial innervation of the EDB muscle. This result was similar to previous studies which reported that ADPN with total innervation of the EDB muscle is a rare condition with the presence of few case reports describing it [11,16,17]. It was postulated that the calculated incidence of total innervation of the EDB muscle by the ADPN is about 0.49% [11].

The ADPN has more than one clinical importance [2,10]. Studying the ADPN can complicate the clinical picture and disturb the interpretation of the electrophysiological studies conducted during the assessment of common peroneal, deep peroneal, and superficial peroneal nerves lesions and injuries, as well as, ADPN neuropathy [1,2,10,13,22–24].

The presence of ADPN with deep peroneal nerve lesion could give the picture of an incomplete deep peroneal nerve lesion instead of a complete one. In this case, there is partial preservation of the function of the EDB muscle in spite of complete loss of function of the tibialis anterior muscle. In this situation, the peroneal CMAP at the ankle stimulation site is unelicited while it is elicited proximally. Needle electromyography (EMG) of the EDB muscle shows evidence of partial nerve lesion [2,3,10,13]. In case of superficial peroneal nerve lesion, the presence of ADPN could give the picture of common peroneal nerve lesion instead of superficial peroneal nerve lesion [2,3,10,13]. In this case, there is weakness of the EDB muscle with weakness of the peroneus longus and brevis muscles in spite of complete preservation of the tibialis anterior muscle. Electrophysiologically, the peroneal CMAP elicited by distal and proximal stimulation sites could be less than normal. Needle EMG of the peroneus longus, peroneus brevis, as well as EDB muscles show an abnormal EMG pattern, while the tibialis anterior muscle shows normal EMG pattern [2,3,10,13]. It is difficult to diagnose cases with partial lesion of the common peroneal nerve with focal conduction block at the transfibular segment having ADPN. The small CMAP obtained at the lateral popliteal fossa stimulation site could be similar to that obtained at the ankle stimulation site. In this situation, the CMAP obtained at below the fibular head site of stimulation could be larger than that obtained in the other two sites of stimulation [19].

Entrapment neuropathy of ADPN could be a cause of chronic ankle pain. The entrapment could be due to compression of the ADPN between peroneus accessories (peroneus quartus) muscle (a common variant muscle of the ankle) and peroneus brevis tendon in the lateral ankle region behind the lateral malleolus [2,15,24,25].

Iatrogenic injury to the ADPN during surgery could be occurred if the surgeon did not put ADPN presence into his consideration [10,15]. This could occur during orthopedic surgery in the ankle region which is usually done through a lateral approach in which the incision is behind the lateral malleolus where the ADPN could be located. Also, this could occur during sural nerve biopsy due to the close relationship between the ADPN and the sural nerve [10,15]. The surgeon should be aware of the presence of ADPN which could be injured during these procedures if it is present [2,10]. ADPN injury could lead to pain in the posterolateral aspect of the ankle region [24]. So, it is recommended to assess the presence of ADPN before doing any surgery in the ankle region and sural nerve biopsy [2,10].

This study had a sampling limitation as it was conducted only in one medical center in one Egyptian governorate. It was difficult to generalize the results of this study to all Egyptian population. Further studies are recommended on a larger scale of Egyptian subjects from different governorates for proper and wider calculation of the prevalence of ADPN among Egyptian population.

**Conclusion**

This study demonstrated that ADPN prevalence in the referred Egyptian sample through electrodiagnostic
studies of lower limbs was 17%, with no sex nor side difference. Recognition of ADPN is essential for proper interpretation of lower limb electrophysiological data, which avoids the error in the diagnosis of common peroneal, deep peroneal, and superficial peroneal nerve lesions.

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Conflicts of interest
There are no conflicts of interest.

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