ABSTRACT
A practical issue in the diagnosis of amyotrophic lateral sclerosis (ALS) is how long the EMG must be observed before a muscle can be declared free of fasciculations with some degree of certainty. To answer this question, the intervals between fasciculation potentials (FPs) were recorded from 53 muscles of 19 ALS patients. The distribution of the FP intervals found across the sample showed that to record a single fasciculation with a probability approaching unity, observation for up to 90 s may be required.

BACKGROUND
Clinical neurophysiologists bear a heavy responsibility in the diagnosis of ALS, there being no reliable biological marker of the disease. Acute and chronic partial denervation seen on EMG provide diagnostic information of lower motor neuron abnormality and its distribution. Although they are found in many other conditions, fasciculations are the hallmark of the condition, and it is difficult to be sure of the diagnosis in their absence. Both denervation and fasciculation potentials (FPs) may be seen in muscles without clinical abnormality emphasising the importance of EMG.

FPs may be recorded by needle or surface electrodes and represent the spontaneous discharge of a motor unit or a part thereof. Needle-recorded FP waveforms can be highly complex, probably due to distal multifocal triggering and intermittent intramuscular axonal block, or may be simple, resembling the waveform of motor units recruited by voluntary action. Their discharge rate is usually described as irregular or random. A recent multi-electrode surface EMG study has shown that intervals may follow either a Poisson distribution, as might be expected from a random rare event, or a more symmetrical distribution of shorter intervals (3–15 ms), generally referred to as double FPs. The former was postulated to be related to spinal motoneuron hyperexcitability, whereas the latter was thought to reflect the phase of axonal superexcitability following the first discharge. Thus, study of FP discharge rate may provide information allowing the neural structure generating the spontaneous potential to be identified.

The electromyographer, when examining a patient with potential ALS, is often faced with the question: how long should I observe this needle recording before I conclude that fasciculations are absent? The question may be less critical if denervation is clearly present or the muscle is weak, but in early cases where the presentation is in a single limb or purely bulbar, the finding of FPs in clinically uninvolved muscles assumes a much greater importance.

This short report attempts to answer this question by providing in a prospective study data on the probability of detecting FPs in relation to observation duration. Many electromyographers would be uncomfortable assigning significance to a single FP and would wait until more were seen to be confident of the finding. Hence, probability values have been calculated for observing one to five fasciculations.

METHODS
FPs were recorded with a concentric needle electrode from 55 muscles (biceps: eight; tibialis anterior: 27; first dorsal interosseous: 17; trapezius: one) in 19 patients with definite ALS according to the El Escorial criteria modified by the Awaji consensus. Patients were selected after EMG had been performed on the above criteria from a sequential group of 61 referred for EMG in whom ALS was a potential diagnosis. Muscles for investigation were selected on clinical grounds with the requirement that muscles from all four limbs and a muscle innervated by a cranial nerve were studied in all patients. An arbitrary criterion was needed as to how long to monitor a muscle for FPs before it was included in the study. If a muscle showed no FPs on EMG after 1 min (the current clinical practice of the author), then that muscle was excluded from analysis. It is acknowledged that this may have excluded some muscles with long interfasciculation intervals but had the virtue of practicality. The duration of disease in the 19 patients was between 3 and 36 months (median: 18 months; IQR: 10.25 months). Recording at a single needle site continued until at least 50 FPs had been recorded. The EMG signal, band-pass-filtered between 5 Hz and 10 kHz, was digitised at a sampling rate of 20 kHz. FPs triggered data collection using Signal 3 software (Cambridge Electronic Design, Cambridge). The time of occurrence of each FP from the start of the recording was logged to the nearest millisecond, and the interfasciculation intervals computed. For each recording, if \( t_1, \ldots, t_n \) represent the times of occurrence of FPs, the intervals \( (t_1 \ldots t_{n-1}) \) were calculated from \( t_{n+1} - t_0, t_{n+2} - t_0, \ldots, t_{n+m} - t_0, t_{n+m+1} - t_{n+m+1} \) and \( t_{n+m+2} - t_{n+m+1} \) for all values of \( n \). The maximum values of each of these series therefore represent the longest interfasciculation intervals in which one, two, three, four or five FPs occurred. The cumulative frequency distribution of \( I_1, \ldots, I_5 \) from the whole data set was then constructed (see figure 1), allowing the probability of 1, 2, 3 or 5 fasciculations occurring with respect to the duration of the recording to be estimated; in
other words, after the needle has been inserted, what range of times may elapse before one, two, three, four or five fasciculations have occurred.

RESULTS
Of the 53 muscles, 16 had normal strength and showed no evidence of acute or chronic partial denervation on qualitative EMG, and eight were severely denervated with fibrillation and marked chronic neurogenic changes. The remainder (29) showed mild neurogenic changes and/or mild weakness. The sample therefore represented the full spectrum of the abnormality encountered in ALS. The duration of recording varied from 51.5 to 776.8 s, giving overall fasciculation rates of from 4.03 to 116.6 FPs/min (first dorsal interosseus: 4.05–67.9 FPs/min, biceps 38.4–116.6 FPs/min, tibialis anterior 4.03–57.1 FPs/min and trapezius 14.6–40.7 FPs/min). The longest interfasciculation interval in the whole data set was 92.2 s. The cumulative frequency distributions for I$_1$ ... I$_5$ are shown in figure 1. From this, it can be seen, for example, that to record five fasciculations with a probability approaching unity, the duration of recording should be 180 s, or the probability of recording 1 FP in 30 s is 0.72, etc. The longest interfasciculation interval has a skewed distribution that could be rendered approximately normal by log transformation. The longest interval found in muscles showing clinical weakness (n=20) had a mean±SD of 23.7±23.0 s compared with that from muscles showing no clinical weakness (n=53) of 25.9±20.0 s. These values are not significantly different (t=0.66, p=0.51, DF=54) using log-transformed data. Similarly, the mean longest interval from upper-limb muscles (n=27) was 21.3±20.8 s, and from lower-limb muscles (n=26) 26.5±21.2 s; again, these are not significantly different (t=1.67, p=0.1, DF=48) using log-transformed data.

DISCUSSION
Secure diagnosis of ALS relies heavily on EMG. It is required to demonstrate lower motoneuron abnormalities in a distribution that cannot be explained by nerve, plexus or root lesions. In early cases where clinical abnormalities are limited to one or two regions, then EMG assumes a greater importance being able to demonstrate EMG changes in clinically unaffected muscles. In limb-onset ALS, it is also particularly important to demonstrate EMG abnormality in muscles innervated by cranial nerves. It remains debatable whether FPs can occur in muscles showing no other EMG abnormality. Some contend that by using quantitative techniques such as macroEMG, FPs occur exclusively in partially denervated muscle. Others using qualitative assessment, as has been used in the present study, find FPs in muscles with no other EMG abnormality. Nevertheless, the confident detection of FPs in ALS is of great practical importance.

The distribution of intervals between FPs in ALS is known to be highly skewed with many short intervals and very few long intervals. The discharge rate may be correlated to the complexity of the FP waveform, suggesting that the degree and recency of collateral reinnervation may be related to discharge rate. At a single needle site, several different FPs may be seen, recognisable by the similarity of waveform components. It would be of interest to study the firing characteristics of individually identified FPs, but the purpose of the present study was a practical one: to determine the longest interval between FPs of any waveform, since these will have the same significance for diagnosis.

The length of time an electromyographer should observe the EMG for fasciculation clearly depends on the clinical weighting to be put on the finding and on the experience of the electromyographer. Practicality is also important; watching EMG for FPs in a perfectly relaxed muscle for more than a few minutes would be difficult for both patient and physician. In early ALS cases where clinical abnormality is limited to one limb, the finding of FPs in the contralateral limb would have greater significance and would therefore warrant a longer search. Any guidance therefore must take into account the clinical situation. However, from these data, it may be suggested that to be certain of recording, say, two fasciculations, the observation should be for 2 min before the muscle can be confidently declared fasciculation-free (figure 1). Similarly, if the muscle shows denervation, and the electromyographer is happy to identify a single fasciculation, the longest time needed to observe the muscle would be 90 s. Of course, in most instances, FPs will be found much more quickly, the above times being the longest needed before deciding that fasciculations are almost certainly absent.

It should also be emphasised that FPs should not be taken in isolation as evidence for ALS; acute and/or chronic denervation in addition in other muscles will be required.

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Detecting fasciculations in amyotrophic lateral sclerosis: duration of observation required
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