Kinematics of individual muscle units in natural contractions measured in vivo using ultrafast ultrasound

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Abstract

Objective. The study of human neuromechanical control at the motor unit (MU) level has predominantly focussed on electrical activity and force generation, whilst the link between these, i.e. the muscle deformation, has not been widely studied. To address this gap, we analysed the kinematics of muscle units in natural contractions. Approach. We combined high-density surface electromyography (HDsEMG) and ultrafast ultrasound (US) recordings, at 1000 frames per second, from the tibialis anterior muscle to measure the motion of the muscular tissue caused by individual MU contractions. The MU discharge times were identified online by decomposition of the HDsEMG and provided as biofeedback to 12 subjects who were instructed to keep the MU active at the minimum discharge rate (9.8 ± 4.7 pulses per second; force less than 10% of the maximum). The series of discharge times were used to identify the velocity maps associated with 51 single muscle unit movements with high spatio-temporal precision, by a novel processing method on the concurrently recorded US images. From the individual MU velocity maps, we estimated the region of movement, the duration of the motion, the contraction time, and the excitation-contraction (E–C) coupling delay.

Main results. Individual muscle unit motions could be reliably identified from the velocity maps in 10 out of 12 subjects. The duration of the motion, total contraction time, and E–C coupling were 17.9 ± 5.3 ms, 56.6 ± 8.4 ms, and 3.8 ± 3.0 ms (n = 390 across ten participants). The experimental measures also provided the first evidence of muscle unit twisting during voluntary contractions and MU territories with distinct split regions.

Significance. The proposed method allows for the study of kinematics of individual MU twitches during natural contractions. The described measurements and characterisations open new avenues for the study of neuromechanics in healthy and pathological conditions.

1. Background

Motor units (MUs) translate synaptic input into muscle motion: the action potentials (APs) discharged by the spinal motor neurons (MNs) in response to synaptic input evoke transient mechanical contractions, called twitches, of the fibres of the muscle unit. Ultimately, global motion is a result of the summation of these muscle unit twitches. The neuromechanics of movement can therefore be studied at the MU level, where the ultimate neural code of movement is translated into mechanical output.

Human neuromechanical control is well explored from both an electrical perspective, providing little spatial detail, and in terms of muscle contraction, mainly treating the muscle as a global force producer. However, how the microstructures in the muscle combine to do so is less clear—the complexity and degree of linearity of the system are unknown. A
representative example of a mechanism relevant for understanding the neuromechanics of muscle control is excitation-contraction coupling (E–C coupling), which is one of the components of the electromechanical delay (EMD)—a metric widely used to study human physiology (Cè et al. 2014, Conchola et al. 2015, Esposito et al. 2016, Smith et al. 2017, Úbeda et al. 2017). E–C coupling refers to the communication between the electrical activity in the skeletal muscle fibre membrane and the release of calcium, eliciting the contraction (Calderón et al. 2014). The ability to measure E–C coupling in vivo in voluntary contractions for individual MUs would provide insights into neural control as the process of modulating MU activities, and would also enable detection and monitoring of pathological conditions in which the E–C coupling is affected, such as myotonic dystrophy 1 (Esposito et al. 2016) and lower back pain (Vasseljen et al. 2006). Coupling time has been measured in electrically stimulated contractions (Nordez et al. 2009, Esposito et al. 2011, Hug et al. 2011a) or in voluntary contractions at the global muscle level with problematic reliance on bipolar electromyography (EMG) (Hug et al. 2011b, Begovic et al. 2014). A method to simultaneously study the electrical and mechanical behaviour of active MUs is required.

Generally, however, methods for MU characterisation can be divided into two broad categories—those which consider the electrical activity, and those which consider the mechanical output. The electrical behaviour of MNs is usually analysed with EMG, either intramuscular (iEMG) or surface (sEMG). Both iEMG and sEMG allow the indirect identification of MN discharges by the analysis of the corresponding muscle unit APs. On the other hand, the mechanical response of entire muscles can be studied using ultrasound (US) (Botter et al. 2013, Dieterich et al. 2017, Tweedell et al. 2019). US has also been used for studying the mechanical response of individual MUs in electrically stimulated contractions (Defieux et al. 2008, Waasdrup et al. 2019, 2021). Furthermore, some studies have suggested that US can even be used at the muscle unit level (Rohlén et al. 2020a, 2020b, Carbonaro et al. 2022). Here, we utilise US alongside a new signal processing pipeline, with minimal processing and assumptions, to extract detailed new characteristics of the kinematics of single muscle units.

The purpose of this study was to provide a proof of concept for the characterisation of individual MU motion for multiple simultaneously active MUs using ultrafast US. We describe and validate the proposed methodology, and we provide results on new physiological measurements not previously possible, including MU-specific E–C coupling time, allowing deeper insight into neuromechanics. We also present the first evidence of single MU twisting during muscle twitches.

2. Methods

2.1. Ethical approval
All procedures and experiments were approved by the Imperial College Research Ethics Committee (reference: 20IC6422) in accordance with the declaration of Helsinki. As the datasets acquired are large, the data were not registered in a public database but may be made available, as appropriate and reasonable, upon request.

2.2. Participants
A total of 12 healthy participants were recruited for this study. Due to synchronisation and system errors, data for 2 of these participants was not used for analysis, leaving ten participants ($n = 10$, 26.2 ± 2.9 yr, 173.2 ± 7.3 cm, 69.1 ± 12.0 kg). Before the experiment, the volunteers were fully briefed on the experiment, provided with a participant information form, and asked to sign an informed consent form.

2.3. Experimental setup and equipment
The method described is based on synchronised acquisition of ultrafast US, high-density surface electromyography (HDsEMG), and force data during repeated sustained contractions performed with feedback on MU activity provided by real-time HDsEMG decomposition. Figure 1(A) summarises how APs generate contractions and shows the information captured by each modality.

2.3.1. Ultrafast US
The US data were acquired with the Vantage Research Ultrasound Platform (Verasonics Vantage 256, Kirkland, WA, USA), equipped with a L11-4 v transducer with 128 elements and centre frequency of 7.24 MHz. Single angle plane wave isonification was used to enable a frame rate of 1000 frames per second with acquisitions of 30 s, resulting in 30 000 frames of US data per recording. The data were beamformed using delay and sum beamforming, resulting in images of 357 pixels axially and 128 laterally, corresponding to pixel dimensions of 0.1 and 0.3 mm. The frame rate and recording duration were chosen as per memory limitations. The values were chosen empirically and proved to be a good compromise between temporal resolution and duration for accurate processing, however variations in these values would also be appropriate as we observed that the method worked over a large range of resolutions and durations. The acquisition and control were performed by custom codes written in MATLAB (Mathworks, Massachusetts, USA).

2.3.2. HDsEMG
The HDsEMG signals were recorded by two electrode arrays of 64 channels (5 columns and 13 rows; gold coated; 8 mm interelectrode distance; OT...
Figure 1. Diagram of the experimental set-up and the underlying human neuromechanical control mechanisms. (A) Motor command is passed from the brain via the spinal cord to the motor neurons (MNs). Each MN innervates a group of muscle fibres. The ensemble of action potential (AP) trains from the pool of MNs innervating a muscle are often referred to as the neural drive to the muscle. Together, the innervated muscle fibres (muscle unit) and the motor neuron constitute a motor unit (MU). APs travel along the MN axon and generate corresponding APs along the muscle fibre membranes. These cause a contraction (shortening and thickening) of the fibres. The ultrasound (US) imaging plane is perpendicular to the direction of the fibres, so a radial expansion is seen in the imaging plane. The velocity mapping algorithm detects axial motion only, so only the components of the expansion perpendicular to the probe are detected. High-density surface EMG (HDsEMG) arrays (for clarity, only one is shown in this diagram) are placed over the surface of the skin, and an online decomposition algorithm is used to estimate the motor unit action potential (MUAP) discharge times. (B) Location of the EMG arrays (rectangles) and location of the US probe (black oblong with dotted red line representing the imaging plane).

Bioelettronica, Torino, Italy). The signals were recorded in monopolar derivation, amplified, sampled at 2048 Hz, A/D converted to 16 bits with gain 150, and digitally bandpass filtered (10–500 Hz), using an EMG pre-amp and a Quattrocento Amplifier (OT Bioelettronica, Torino, Italy).

2.3.3. Force
An ankle dynamometer was used to measure the dorsiflexion force. The leg and foot were constrained in the dynamometer using foam padding and secured using straps to provide stability and to eliminate other motions, keeping the foot at an angle of 90° to the leg. The force data were fed through a Forza force amplifier (OT Bioelettronica, Torino, Italy) directly into the auxiliary port of the Quattrocento amplifier.

2.3.4. Synchronisation
A 1 μs active low output was fed from the Verasonics US system into an Arduino UNO that in turn outputs a 490 μs pulse, elongated to allow for detection at the 2048 Hz sampling frequency of the EMG amplifier. The trigger was recorded at the start and end of each recording period and used to align the data in post-processing. To test the accuracy of synchronisation and alignment of the US series and the HDsEMG signal, the HDsEMG grid was sharply perturbed such that a peak was seen in the electrical (as artefact) and velocity signal obtained from the US series. These peaks were seen to align within the expected error of ±0.5 ms resulting from the US frame rate, confirming the validity of the synchronisation method.

2.3.5. Setup
The experiment was conducted on the tibialis anterior (TA) muscle. The motivation for using the TA in this study was threefold: it is a large, long muscle allowing space for all the sensors used; it is superficial, resulting in high-quality HDsEMG decomposition (Del Vecchio et al 2020); and its fibres are long and have a relatively low pennation angle (Sopher et al 2017) (see figure 2). The latter property allowed for mounting the US probe and the HDsEMG arrays in separate areas whilst ensuring they still overlay the same fibres, and ensuring more uniform velocity fields within the imaging plane. In all participants, the experiment was performed on the (dominant) right leg.

Before beginning the experiment, the skin over the TA was shaved, lightly treated with a chemical abrasive, and cleansed using an alcohol spray. The muscle belly was then identified by palpation whilst the participant was guided to contract and relax the TA. A HDsEMG electrode array was placed over the proximal part of the muscle, parallel and lateral to the tibia. The grid was positioned such that all electrodes were on top of the TA belly. A 1.5 cm gap was left
distally with respect to the first array and a second array was placed over the distal part of the muscle, also parallel and lateral to the tibia. In some participants, the distal electrodes of the second array fell outside the TA belly, which resulted in small signals and did not impact the results. The electrodes were placed along the estimated direction of the muscle fibres and secured using Tegaderm Film Dressings and medical self-adhesive bandages.

Next, the US probe was placed in the gap between the EMG electrodes, with the imaging plane perpendicular to the length of the muscle fibres (see figure 1(B)). The innervation zone of the TA has been found to be at 33.8 ± 7.1% of the distance between the length of the muscle (Beretta Piccoli et al 2014), hence it is likely to underlie the more proximal EMG grid. However, large inter-subject variability has been seen in innervation zone location (Bowden and McNulty 2012), hence the use of two grids. A water-based US gel was used to improve the coupling between the probe and the skin. The US probe was held by a custom-built probe holder and secured in position using straps. The participant sat in a chair in a comfortable position with the leg strapped into the ankle dynamometer. The US system was turned on to perform real-time imaging and small adjustments were made to ensure the TA muscle belly was covered. Finally, a computer screen was adjusted to a comfortable distance and position to provide visual feedback during the experiment. The described configuration of separate locations for the EMG grids and US probe was preferred over the possibility of using grids transparent to US directly under the US probe (Botter et al 2013) since transparent grids have not been tested for single muscle unit motions (see also section 4).

2.4. Experiments

Real-time online decomposition of the HDsEMG signals into MU discharge times was performed using the methodology of Barsakcioglu et al (2021). The identified spike trains were provided as visual feedback to the participant throughout the experiment. This approach ensured continuous activation of the target MUs, and allowed the participant to precisely control the number of recruited MUs.

The online decomposition was trained on a contraction at 10% maximum voluntary contraction (MVC), maintained for 41 s. Once the online decomposition model was trained, the subject was asked to slowly increase the dorsiflexion force whilst the experimenter inspected the decomposition to determine whether the model was satisfactorily trained, and the noise level was acceptable. Model training was repeated when deemed necessary (this was never done more than once during the experiments).

With the decomposition trained, the visual feedback displayed the MU discharge times in real-time. The subject was asked to slowly increase the dorsiflexion force, thereby slowly increasing the number of recruited MUs displayed on the screen. The order of recruitment of the identified MUs was noted. The subject was then trained to recruit individual MUs and to incrementally increase the number of MUs in a controlled manner. Providing visual feedback was a vital part of the training as fine recruitment of low threshold MUs is a complex task.

Once the participant was comfortable with this control method, the recordings commenced. The subject was instructed to recruit a single MU and to keep it active for 30 s whilst the US and HDsEMG recordings were taken. The subject then rested for 60 s before repeating this process for the same MU two
Table 1. Participant and recording information. ‘MUs detected’ is the total number of motor units (MUs) that were recorded using EMG at any time during an ultrasound (US) recording. ‘Quality MUs’ is the total number of MUs that were properly visible in the processed US recordings and of sufficient quality for identification. ‘Total Paired recordings’ is the total number of paired recordings made for each subject. ‘Quality Paired Recordings’ is the total number of recordings with at least one MU of sufficient quality i.e. the number of recordings used in data analysis. ‘MU-STA curves obtained’ is the total number of motor unit spike-triggered average (MU-STA) curves produced for each participant.

| Participant | MUs Detected | Quality MUs | Total Paired Recordings | Quality Paired Recordings | MU-STA curves obtained |
|-------------|--------------|-------------|-------------------------|---------------------------|------------------------|
| 1           | 8            | 6           | 18                      | 12                        | 33                     |
| 2           | —            | —           | —                       | —                         | —                      |
| 3           | 7            | 7           | 19                      | 14                        | 50                     |
| 4           | 5            | 3           | 18                      | 12                        | 19                     |
| 5           | 6            | 5           | 19                      | 17                        | 51                     |
| 6           | —            | —           | —                       | —                         | —                      |
| 7           | 4            | 4           | 14                      | 12                        | 31                     |
| 8           | 7            | 5           | 18                      | 17                        | 39                     |
| 9           | 7            | 5           | 18                      | 14                        | 40                     |
| 10          | 7            | 4           | 18                      | 18                        | 28                     |
| 11          | 9            | 6           | 17                      | 17                        | 57                     |
| 12          | 8            | 6           | 20                      | 15                        | 42                     |
| Total       | 68           | 51          | 179                     | 148                       | 390                    |

Further times. Each 30 s paired recording of HDsEMG and ultrafast US was considered a single trial. The three trials were then repeated for up to six different MUs. If fewer than 6 MUs were identified, the process was only performed up to the maximum number of identified MUs.

2.5. Data analysis and processing

2.5.1. Datasets

For one subject the synchronisation signal failed, and for another the HDsEMG decomposition failed, resulting in valid data from ten subjects. For an example participant (participant 1), 8 MUs were detected by HDsEMG and 18 paired recordings of HDsEMG and US were taken. Of these 8 MUs, six were in the correct plane to be seen by the US probe, resulting in 12 usable paired recordings. In each of these 12 recordings multiple MUs were active, and the number of active MUs varied. As such, from these 12 recordings, 33 MU-spike-triggered averages (STAs) were computed. Overall across 12 participants 51 MUs were identified in both US and HDsEMG and 390 MU-STA curves were obtained. For each participant the details are summarised in Table 1.

2.5.2. Data processing

From the online decomposition, the discharge times of the identified MUs were available. However, each discharge was identified with some variability with respect to the AP waveform. Since in the current application it was important to define the precise onset of the MU discharge, we estimated the required temporal shift so all APs would be aligned in time. The shift was identified by means of an STA of the HDsEMG signals over each channel, producing the average AP shape for each MU. A double differential AP shape was calculated and the channel with highest signal-to-noise ratio was selected, and the onset of the AP was defined as the first time sample with amplitude higher than five standard deviations of the baseline signal (Ibáñez et al 2021). The time difference between the onset and the centre of the STA window was the shift by which all AP spikes of the MU were shifted, such that the corrected AP discharge time reflected the onset of electrical potential at the neuromuscular junction in a consistent manner for all discharges.

For the ultrafast US data, each trial comprised 30 000 frames (357 × 128 pixels each) that were used to generate a series of velocity maps using 2D autocorrelation velocity tracking (Loupas et al 1995) with a sliding window of two frames, corresponding to 2 ms. This resulted in 29 998 frames (357 × 128 pixels) with bipolar amplitude representing axial velocity (negative pointing away from the probe). The recordings were then further low-pass filtered (0.5 MHz) along the axial direction and band-pass filtered (5–100 Hz) along the temporal direction, removing noise from outside the expected temporal range of the response.

Next, a STA of the US velocity maps was performed using the discharge times identified from the HDsEMG decomposition. For each identified MU, a ±50 ms window (100 frames in total) of velocity data around each discharge was selected (figure 3). All windows were averaged, resulting in 100 frames of STA US maps (US-STA maps) for each MU (figure 4(A)).

Supplementary video 1 in the supporting information shows an example of the 100 frame US-STA i.e. the average muscular movement caused by an individual MU discharge. In this representative video, a large part of the movement caused by the target MU is hidden by the motion of non-muscular structures. These high noise areas need to be removed for
Figure 3. Diagram of the spike-triggered average (STA) method employed. For each motor unit (MU) and identified discharge, a window around the discharge timing is selected. All selected windows are summed and averaged, resulting in a single STA window for each MU. Note that this process is performed on the velocity fields obtained from the ultrasound image series.

Figure 4. Process to highlight motor unit (MU) related motion. For each MU a window of 50 frames before and 50 frames after each action potential (AP) time is selected (shown in panel (A), where $M = 100$ frames, $N = $ total number of APs in that recording, $X =$ width of the image, and $Y =$ depth of the image) and a spike-triggered average (STA) technique is employed to obtain the STA and the variance of the STA, as shown in panel (B). The STA is divided by its variance and summed over time to obtain a single map of intensities, where high intensity represents areas of high motion and low variability in the STA. The values are also multiplied by a direction factor (as shown in panel (C)) which separates segments of the STA map moving in opposite directions in response to the AP.
an accurate analysis of the muscle unit movement. For this purpose, for each US-STA window, the variance across all windows was calculated (figure 4(B)). To determine the direction of motion of a pixel as induced by the MU activity (either positive—towards the probe—or negative—away from the probe), we performed a subtraction of the sum of the second half of the STA from the first (figure 4(C)). For each pixel the sum over the squared STA divided by the variance (with the convention of a positive sign for upwards and negative sign for downwards motions) was calculated, highlighting regions of intense (high amplitude) and consistent (low variance) movement, whilst penalising regions of non-consistent movement (e.g. tibia movement). Lastly, the map was downsampled axially by a factor of 3 (119 × 128 pixels wide; 0.3 × 0.3 mm per pixel) and the regions of no interest (the rows after the maximum sEMG range) were removed, leaving maps of size 39 × 120 pixels (11.5 × 38.4 mm). Each of the final maps, which we will term MU activity maps (MUAMs), represents the muscular movement caused by an individual MU discharge during a voluntary isometric contraction. In summary, the recorded images have approximate width 38 mm and depth 35 mm. These images are cut to remove regions from which no EMG activity can be detected, resulting in images with approximate width 38 mm and depth 12 mm. The pixel size of these final images is 0.3 × 0.3 mm. As such, the resolution is not high enough to detect single muscle fibre movement as fibres in the TA have cross sectional area of approximately 5000 µm² (Wåhlin Larsson et al 2008). However, with this resolution, muscle units, whose approximate width in the TA is approximately 5 mm (Diószeghy 2002), can be resolved.

2.5.3. Muscle unit area of motion identification
The MUAMs show, for each pixel, the effect caused by the discharge of a specific MU, with high values (positive or negative) representing regions of large motion that are highly synchronised with the MU discharge, whilst low values represent regions with either small or highly desynchronised motion. In some situations, an MUAM may show separate regions of positive and negative velocity associated with a single muscle unit motion. In these cases, the two regions were treated separately but assigned to be part of the same MU. To properly define which pixels were associated to the movement of a muscle unit, a thresholding approach was used. For each MUAM, pixels with absolute value greater than 65% (chosen empirically) of the maximum amplitude were considered as part of the region of movement due to the MU activity. We will refer to the pixels constituting this region as the MU motion domain. In contrast to the MU territory, defined as the cross-sectional area containing the muscle fibres of an MU, the MU motion domain is therefore defined as the region of muscle tissue that shows movement with amplitude and synchronicity over a predefined threshold when an MU discharge occurs. Whilst the MU territory refers to the MU anatomy, the MU motion domain refers to the mechanics of the muscle unit.

2.5.4. MU mechanical twitch profile
The MUAMs were used with the US-STAs to generate an STA of the mechanical twitch of the MU (MU-STA curves). These were calculated by averaging the US-STA pixels curves inside the MU motion domain. One MU-STA curve twitch profile was generated for each MU (two if the MU caused both a positive and negative movement in separate regions) in each trial. In this work, twitches refer to the movement of the fibres in terms of a displacement velocity.

Each extracted MU-STA curve is a unique twitch profile that was characterised the parameters shown in figure 5(A). Limited work in this area has resulted in unclear and inconsistent naming of parameters and, as such, our nomenclature may not be in agreement with other works. One extracted feature was the duration of the mechanical twitch—the time over which the tissue drastically changes its velocity profile. This corresponds to the time interval over which the tissue accelerates, thus the boundaries of this interval are inflection points. In some cases (figure 5(B)) the onset of the mechanical twitch can be easily identified by the inflection point of the velocity, however in other cases (figure 5(C)) the twitch occurs during a later stage of relaxation of the previous twitch, so although there is a sharp velocity change, the velocity inflection point cannot be used to define twitch onset. As such, the peak in the second derivative was used to mark the onset of the mechanical twitch.

Another feature of interest was the contraction time, defined as active or total. The active contraction time was defined as the interval between velocity zero crossing and the inflection point. The total contraction time starts at the same point, however it ends at the second zero crossing before the next spike. Although both features represent the time where the tissue is in the ‘contracting state’, the active contraction time accounts for the time over which the twitch is actively accelerating the tissue, whereas the total contraction time includes the full duration of the ‘contracting state’—the total contraction time includes both the time when the fibres are actively generating the contraction, as well as the time when the acceleration is due to the elastic recovery forces.

Finally, we also calculated the activation delay, defined here as the time between the onset of the electrical AP and the onset of the mechanical twitch. Currently, the EMD is measured as the delay between the onset of the detected electrical activity and the onset of force production, and often its measure has significant inaccuracies due to the use of bipolar EMG at a distance from the motor point (Cavanagh and Komi 1979). As such, in literature, the EMD includes
Figure 5. Measurements extracted from the motor unit spike-triggered average (MU-STA) curves. (A) The dotted AP line is the time of the onset of the electrical action potential (AP) measured using high-density surface EMG. Note that before the effects of the next twitch begin, the fibre is still relaxing from the previous twitch, thus the velocity is non-zero and changing prior to the AP. Following the AP, the velocity reverses due to the twitch, and the time between the AP and the turning point is the activation delay. The fibre is then accelerating due to the twitch between the first and second inflection points—we refer to this as the duration of the mechanical twitch. Between the two zero crossings, the fibre is in a contracted state—thus we break this region down into active contraction time, where the fibre is being actively contracted, and total contraction time, which also accounts for the relaxation back to the neutral position. (B) An example velocity profile for a motor unit (MU) where the interspike interval is low, so an inflection point is seen at the onset of the mechanical twitch. (C) An example velocity profile for a MU where the interspike interval is high, so the previous mechanical twitch is in a later stage of relaxation resulting in no turning point at the onset of the twitch. To enable twitch onset detection in both cases, the peak of the second derivative is used (solid black line). In each case, the dotted black line is the firing time from the EMG, highlighted by a red arrow above the plots.

the time lag due to the synaptic transmission, the E–C coupling, the muscle force transmission on the series elastic components, the aponeurosis, and force transmission on the tendon. Our technique allows for the measurement of the delay between the onset of voluntary motor unit action potentials (MUAPs) and the first measurable acceleration on the fascicles of single muscle units. Therefore, this measure includes the time lag caused by the E–C coupling and force transmission on the series elastic components (between the innervation point and the US measurement plane). However, as the force transmission travels at $\sim 30 \text{ m s}^{-1}$ (Morimoto and Takemori 2007, Nordez et al 2009) and the end-plates for the detected MUs were within 4 cm distance to the US plane, the resulting delays were less than 1 ms. Thus, the measured activation delay mainly represented the E–C coupling, making our measurements the first to report individual MU E–C coupling delays.

2.5.5. Processed dataset

Following the process described above, all 179 paired recordings were processed. However, 31 recordings were discarded as the HDsEMG decomposed spike trains were poorly identified. A total of 68 MUs were identified across all participants in the HDsEMG recordings, 51 of which (75%) could also be detected in the US recordings. This discrepancy is due to out of plane MU activity, a problem which is further explored in the Discussion. As such, 390 MU motion domains and mechanical twitch profiles from 51 MUs from ten participants.
2.5.6. Statistical analysis
The data presented in this paper are intended to act as a proof of concept, showing what the set-up and processing pipeline can achieve. For each value and graph, the number of data points, \( n \), is stated in the accompanying text. For all histograms and box plots \( n = 390 \).

3. Results

Figure 6 shows an example of the two signal modalities (EMG and US) plotted together, as well as their STAs. For each MU activation, the velocity inside the MU motion domain changes in a synchronous manner. Figure 7 shows an example of the outputs for a trial with four identified MUs. The identified units have variable discharge rates (figure 7(A)), as is expected when the force level is close to the recruitment threshold of the MU. The mean firing rate of the MUs was 9.8 ± 4.7 Hz. Figures 7(B) and (C) compare the shapes of the electrical STA and the MU-STA—the electrical STA is unique to a given MU while the MU-STA are similar in this example. The MUAMs in figure 7(C) are truncated at approximately 10 mm as throughout the experiment no MUs were detected below this level, because of the relatively superficial detection volume of the surface EMG (Merletti and Muceli 2019).

3.1. MU spatial motion domain
To analyse the consistency of the area assigned as the MU motion domain across repeated trials, the percentage overlap of the spatial domains for each MU across different trials was calculated. The results are shown in figure 8 for each participant, and the average for all participants was 47.0 ± 17.4%. In contrast, the percentage overlap between a given MU motion domain and the motion domains of other MUs was 6.9 ± 6.8%. This showed that MU motion domains were more similar to the motion domains of the same MU in different trials than to those of other MUs for all participants. Although a 47% of overlap across trials may seem a low value, it incorporates error resulting from thresholding, as well as morphing and shifting due to other active MUs. The consistency in location of MU domain can be seen for representative MUs in figure 9.

3.2. Mechanical twitch reproducibility
Figure 9 shows, for two representative MUs, both the MUAMs (left panel) and the normalised MU-STA curves (right panel) obtained for different trials (Example MU 1: 17 trials, Example MU 2: 8 trials). Each grey curve is the MU-STA velocity profile, averaged over the MU motion domain. These curves show high consistency across trials, despite changes in conditions (differing number of active MUs). For all 390 MU-STA curves, the average standard deviations between curves from the same unit was 6.69%.

From these results, we concluded that the MU-STA curves were highly repeatable across force levels. Whilst it is clear that the MU motion domain morphs and shifts at differing force levels, the twitch profile is consistent across force levels, suggesting higher linearity in the temporal domain than in the spatial domain.

3.3. Mechanical twitch duration, contraction time, and activation delay

Figure 10(A) shows the histogram of activation delays, with average value 3.8 ± 3.0 ms. A small proportion (3.8% of 390) of calculated activation delays were smaller than zero, suggesting an onset of mechanical motion prior to the electrical AP, due to numerical errors resulting from noise. Figure 10(B) shows a histogram of the mechanical twitch duration, with average value 17.9 ± 5.3 ms. Figures 10(C) and (D) show histograms for contraction times, with average values 12.1 ± 4.0 ms and 56.6 ± 8.4 ms for the active and total contraction times, respectively. All these measures are shown here for the first time for individual MUs in voluntary contractions.

3.4. Muscle twisting

In some instances, we observed regions of both positive and negative motion, aligned depth-wise, in response to AP discharges. On average, 17% of the MUs identified in the experiment across all participants exhibited this behaviour. These can be explained by a twisting of the muscle unit, where one group of muscle fibres moves towards the probe next to a group of fibres moving away from the probe. An example is shown in figure 11(B). This is the first observation of this phenomenon in natural contractions, and is in agreement with the twisting seen in electro-stimulated contractions (Deffieux et al. 2008). In the aforementioned study a fibre bundle with an unknown number of recruited MUs was activated by stimulation, thus twisting could not be reliably accredited to an individual MU. However here, single MUs are studied, and fibre bundle twisting is seen. As such, our results suggest that twisting can occur on an individual MU level. The source of this twisting is unclear and could be attributed to tendon attachment, to the surrounding connective tissue, or could be a fundamental contractile property of the MUs.

4. Discussion

In this study we have proposed a method suitable for the kinematic study of individual MU twitch properties, with specific focus on extraction of E–C coupling time. This provides new perspectives in physiology for the study of muscular contractions at a granular level in relation to neuromechanical control. Furthermore, the approach could open up new avenues for the study and monitoring of pathologies that relate to muscle mechanics, and those which relate to the translation of motion from the electrical to the
Figure 6. Synchronisation of velocity and EMG detected firing times. (A) Example of average velocity inside motor unit (MU) motion domain for an identified MU (orange line), and action potential (AP) discharge times from EMG decomposition (black dotted lines) showing synchronicity. This is a 1 s interval extracted from a 30 s recording. (B) Motor unit action potential (MUAP) found by means of spike-triggered average (STA) on the EMG signal (black line) and motor unit spike-triggered average (MU-STA) curve for the same MU (orange line). The black dotted line is the corrected firing time—the time at which the double differential of the highest signal to noise channel of the EMG array is above five standard deviations of the baseline noise level. The MU-STA is relaxing from the previous twitch hence the velocity is changing prior to the discharge time.

Figure 7. Example set of outcomes for a recording. (A) Summed raw EMG signal (top) across the 64 recording electrodes in the high-density surface EMG array and the action potential (AP) discharge times of four of the identified motor units (MUs)—each of the units is given a label as shown to the left of the plot. A 10 s excerpt of the 30 s recording is shown. (B) Motor unit action potentials (MUAPs) for each of the identified MUs prior to correction to align activity onset, found by performing STA on the EMG signal. (C) Motor unit activity maps (MUAMs) (left) and motor unit spike-triggered average (MU-STA) curves (right) for each of the four MUs. The MUAMs show the MU motion domains, and the MU-STAs show the average velocity in these regions in response to an AP. For all STA plots, the AP occurs at 0 ms.

mechanical domains. Many myopathies cause either shortened or elongated E–C coupling times, resulting respectively in a loss of function or a gain in function, which in turn results in hypersensitivity (Marty and Fauré 2016). By using this coupling time as a marker, progression of myopathies as well as treatment responses can be monitored over time using our proposed technique, without need for invasive measurements. Within this work, we have adapted the commonly used STA technique by adding a pixel-wise
variability weighting to penalise inconsistency across spike occurrences, and thus minimise the contributions from pixels with large motion not associated with the spikes. This has implications for other applications where relevant signals may be masked, or where long recording times are not possible.

Current methods to study MU motion within a muscle suffer from several limitations. Often, these use electrical stimulation to elicit motion (Deffieux et al. 2008), evoking compound contractions rather than individual MU activity and resulting in contractions different to natural motor control. Further limitations include the use of systems with low temporal resolution compared to the durations of the involved physiological processes (Botter et al. 2013), low spatial resolution to properly separate contributions from multiple MUs (Birkbeck et al. 2020), or the deployment of invasive techniques, impacting the mechanics of the system and enabling the detection of just one MU at a time (Rohlén et al. 2020a). The use of ultrafast US to decompose contributions to muscle motion from individual MUs has been previously attempted using spatio-temporal independent component analysis (Rohlén et al. 2020a, 2020b, Carbonaro et al. 2022), whilst we propose an approach which isolates these contributions by directly using the known MU discharge times.

A method that shares some common characteristics with US is mechanomyography (MMG). This is a technique that detects the deformation of the surface of the skin overlying a muscle, as due to the motion of the muscle units (Orizio 1993, Bichler 2000, Kaczmarek et al. 2005, 2009, Cescon et al. 2008). However, whilst MMG detects motion at the surface of the muscle, thus distant from the contracting muscle units, US can detect local muscle motion within the muscle unit cross section. How intramuscular motions of individual MUs combine to deform the muscle surface, and hence how they contribute to the generation of the MMG signal, is complex to study and unclear as of yet (Orizio et al. 1996). As such, by using the current approach to study muscular deformation at the single MU level and integrating additional MMG measurements, it may be possible to create an accurate biophysical model for the propagation of mechanical waves due to MU motion to the muscle surface, hence providing a method to accurately develop MMG generation models.

4.1. Technique validation

The online decomposition algorithm for the HDsEMG has been validated elsewhere, by comparison both with offline decomposition algorithms and with state-of-the-art iEMG techniques (Barsakcioglu et al. 2021). The rate of agreement between this online decomposition and validated offline decomposition algorithms was greater than 90%. Although offline decomposition has slightly lower accuracy than online decomposition, it has been used successfully on the TA in control tasks (Bräcklein et al. 2022) with
similar conditions to the current study. As such, by using these decomposition results to process the velocity maps, we ensure that our STA process identifies MU activity from a single unit. Furthermore, our STA processing involves dividing the velocity maps by the variance across each spike time, hence if no consistent movement was present the processed maps would be uniformly zero. The processing ensures that regions we are extracting are specific to that unit; the use of the division by the variance in the STA process is itself a validation that the extracted information is related to the single triggering units since otherwise the STA normalised by variance would tend to a null response.

Further, we validated the technique as a whole by analysing the repeatability of the MU-STA twitch profiles, and by comparing the MU motion domains across trials. The low standard deviations of the MU-STA curves found for the MUs across different trials showed the repeatability of the method, despite the in vivo nature of the work, and the variation in force level across different trials.

Spatially, MU motion domains were found to have higher similarity to the motion domains of the same MU in different trials than to that of different, simultaneously active, MUs (on average, 47% vs 7% respectively). The former shows high spatial consistency of MU motion domain location serving as evidence that we are correctly identifying consistent MU motion domains. As expected, this metric of consistency resulted in values lower than 100% due to factors such as probe and muscle shifting, translation and muscle morphing due to other active MUs, concomitant activation of different MUs, noise, and thresholding effects.

Although these values show that the method successfully identifies and separates contributions to muscular motion from multiple simultaneous MUs, they also highlight that even constant isometric contractions imply complex dynamics at the muscle unit level, resulting from mechanical coupling with neighbouring units. Further evidence for the validity of the technique is the agreement of some of the measured physiological parameters with the
corresponding parameters identified with different approaches in previous studies, as discussed in the following section.

4.2. Physiology of the muscle unit

The mean firing rate of the MUs was 9.8 Hz, close to the lower limit for voluntary tasks (Fuglevand et al 1993). This was made possible by the use of real-time MU firing feedback, allowing for participants to ensure they maintained a low MU discharge rate. Given that the interspike interval is known to affect the estimates of MU force twitches when using spike-triggered averaging methods (Thomas et al 1990, Kossev et al 1994), we chose to study units at low firing rates to minimise the effect of subsequent twitches and for consistency. Further studies could focus on looking at the changes in the measured physiological parameters with changes in discharge rate.

It should be acknowledged that, due to the pennation angle of the muscle fibres in the TA, the cross-sectional images of the muscle do not correspond to the physiological cross-sectional area. As such, a bulging circular region of fibres is not expected to be seen. Nonetheless, the influence of this factor is presumably small because of the small pennation angle, as can be observed in figure 2. The actual region of motion is more complicated, due to the angle of the fibres as well as the compressibility of surrounding tissue and the effects of the pressure from the attached probe. However, this motion is still synchronised with the electrical events, and localised to a region.

![Figure 10](image)

Figure 10. Histograms of temporal features extracted from the velocity profiles of motor unit twitches. (A) Histogram of activation delay. A small proportion (3.9%) of results were deemed to be non-physiological as these suggest motion of the MU prior to electrical activity. These have been left unshaded in the histogram to highlight that they are attributed to noise. (B) Histogram of duration of mechanical twitch. (C) Histogram of total contraction times. (D) Histogram of active contraction times. For all histograms, \( n = 390 \) from 51 MUs across ten participants.

Our analysis in the temporal motion domain yielded three mechanical twitch defining parameters: twitch duration, active contraction time, and total contraction time. In their work, Rohlén et al identified a parameter called ‘twitch duration’ (approximately 50 ms) (Rohlén et al 2020b) and Deffieux et al identified a parameter called ‘contraction time’ (70.1 ± 2.0 ms) (Deffieux et al 2008), analogous to our ‘total contraction time’ (57.2 ± 8.7 ms). Our results are within one standard deviation of those by Rohlén et al (2020b). Differences among studies may be due to the different muscles investigated, and methodological differences. On the other hand, larger differences in comparison to the results by Deffieux et al (2008) are likely due to their use of electrical stimulation.

From the time-domain analysis, we extracted the activation delay, i.e. the time between the electrical AP onset and the onset of motion caused by the
twitch. We believe that the main contribution to this value is the E–C coupling, and this work therefore represents the first attempt to measure the duration of the E–C coupling in individual MUs in sustained voluntary contractions, which has potential impact for further study of neuromyopathies, myopathies and other conditions known to affect certain aspects of AP transmission (Orizio et al. 1997, Granata et al. 2000, Kaneko et al. 2002). In electrically stimulated contractions, the time between the stimulation and the onset of fascicle motion is attributed to the combination of E–C coupling and synaptic transmission (6.05 ± 0.64 ms (Nordez et al. 2009), 5.57 ± 1.37 ms (Hug et al. 2011a)). Whilst these studies measure the time between electrical stimulation and fascicle motion, our study instead measures the time between onset of electrical activity as detected by HDS-EMG and onset of fascicle motion, thereby not including the time for synaptic transmission. We would therefore expect smaller latencies in our study as it accounts for E–C coupling only. Another stimulation-based study instead measured the time between EMG onset and fascicle motion (2.2 ± 0.3 ms (Esposito et al. 2011)), which therefore only accounts for E–C coupling, and is closer to our value of 3.8 ± 3.0 ms. It must be noted that these measurements of the activation delay require a very accurate synchronisation between electrical and mechanical measures as the delays are very small. For example, in previous work, the electrical activity seems to follow rather than precede the onset of mechanical activity (see figure 3 in Rohlén et al. 2020a), which is likely due to issues in synchronisation or processing.

It is known that the EMD is more than two times greater for voluntary contractions than stimulated contractions, which is often attributed to the reversed recruitment order resulting in faster force development during stimulated contractions (Hopkins et al. 2007). However, some studies also suggest that the E–C coupling is also greater in voluntary contractions than stimulated contractions (19.21 ± 6.79 ms (Begovic et al. 2014)), in contrast to our results which well align with those reported for stimulation studies. This may be explained by the use of bipolar EMG in these previous studies and by the influence of electrode location with respect to the innervation zones. For example, for electrodes placed at a distance of 4 cm from the innervation zone, there would be a detection delay due to propagation along the muscle fibres of approximately 10 ms (Hug et al. 2011b). Furthermore, the use of A-mode US to measure the muscle thickness and estimate the onset of fascicle motion may introduce errors due to muscle morphing and A-mode placement. This was circumvented in the current study by using B-mode US.
with a processing pipeline to identify individual MU motions. Our results suggest that E–C coupling is similar in stimulated and voluntary contractions.

Spatially, we defined a ‘MU motion domain’ as the region of muscle which moves with magnitude and synchronicity above a defined threshold when an AP travels along the muscle unit. Anatomically, MU territories have been defined as the cross-sectional area of the innervated fibres by a MU. MU territories have been studied using glycogen depletion experiments (Edström and Kugelberg 1968, Garnett et al 1979, Bodine et al 1988) and scanning EMG (Stålberg and Dioszeghy 1991). The latter method analyses the MU territory in one dimension, thus the measured size depends on the location of the needle intersection with respect to the MU cross-section (Gootzen 1990). Furthermore, ‘silent zones’ (regions of no EMG signal) are seen in these measurements, which may be a result of the innervation being scattered in different portions of the muscle (Stålberg and Dioszeghy 1991). More recently, MRI was paired with electrostimulation to measure human MU territories in vivo (Birkbeck et al 2020). These studies found MU regions which they categorised as either circular, elliptical, crescent shape, spider, or split, corresponding well with our MU motion domain shape, which varied largely between MUs (figure 11(A)). The MRI experiments did not detect overlapping MUs due to methodology constraints, whereas we identified overlapping MU motion domains in every participant. On the other hand, split territories were not identified in a study using ultrafast US (Carbonaro et al 2022), presumably because of methodological constraints. Our work therefore presents the first method able to detect both split and overlapping regions of activity.

4.3. Limitations and future applications

While the work presented marks a novel approach to the study of MUs, there are some limitations of note. The results presented only show the technique working on low-force contractions (maximum 10% MVC, resulting in recruitment of slow MUs with lower recruitment thresholds). Translating the results to higher MVCs is not straightforward as they generate high global motion that obscures the individual MU discharges. Higher forces also increase muscle stiffness and increase the likelihood of probe movement and slippage. So far, the processing assumes that the probe remains stationary throughout the recording, which is a valid assumption for low forces; however, for higher forces (or dynamic contractions) this would not hold and considerations for its movement would have to be made.

Due to the use of HDsEMG to identify MUAPs, our methodology currently can only detect MUs near the surface of the skin. However, had we used iEMG we would detect a smaller number of individual units within close range of the needle tip (Lowery et al 2006), and the needle (for the case of needle recordings) would cause mechanical coupling disrupting the results. In principle this can be solved by directly decomposing the US in MU discharges (Ali et al 2020, Rohlén et al 2020a, 2020b), however further validation on the basic assumptions underlying the mechanical components and coupling of MU twitches is necessary.

On the processing side, the autocorrelation velocity tracking algorithm employed is limited to one dimension, however the motion generated by the fibres contractions is multidimensional. Expanding this into a two-dimensional tracking so both forms of the fibre cross-sectional expansion movement can be tracked could be considered. On the same note, the resolution of our US image is limited so individual muscle fibres cannot be detected.

Finally, a formal method of selecting the threshold for producing the MUAMs could be implemented by, for example, optimising the threshold by comparison with the dimensions of MU as measured using scanning EMG. In another study, the spatial representation of the MUs in the biceps brachii appear to span a large portion of the muscle region (see figure 4 in Carbonaro et al 2022), in contrast to our narrower domains. This discrepancy may be a result of the applied methodology in this previous study since EMG was used as a constraint for selecting the region, or of our use of an empirically chosen thresholding value.

5. Conclusion

In conclusion, we have utilised synchronised HDsEMG with online MU decomposition and ultrafast US to isolate contributions to muscle motion from individual MU twitches. In doing so, we have been able to measure the E–C coupling time under non-stimulated conditions for individual MUs. This methodology and the results it yields open up avenues for monitoring pathology development and treatment, and could allow for further study of other features of the MU.

Data availability statement

As the datasets acquired are large, the data were not registered in a public database.

The data that support the findings of this study are available upon reasonable request from the authors.

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Figure 1 was created using BioRendr.com.

Authors’ contribution

All experiments were performed at the Department of Bioengineering at Imperial college London. E L: developed concept, wrote and developed code for
control of Verasonics system, for synchronisation, and for post-processing and data analysis, designed custom probe holder, did in vivo experiments, analysed data, wrote paper. B S: developed concept, did in vivo experiments, analysed data, wrote paper. D B: developed concept, developed code for online EMG decomposition as well as closed-loop biofeedback framework and graphical user interface. D F, M T, J I P: developed concept. All authors revised the manuscript and provided intellectual feedback. All authors have read and approved the final version of this manuscript and agree to be accountable for all aspects of the work in ensuring that questions related to accuracy or integrity of any part of the work are appropriately investigated and resolved. All persons designated as authors qualify for authorship, and all those who qualify for authorship are listed.

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Conflict of interest

None to declare.

Consent for publication

Not applicable.

Ethics approval and consent to participate

All participants signed informed consent before participation in the study. Procedures and experiments were approved by the Imperial College Research Ethics Committee (ICREC reference: 20IC6422) in accordance with the declaration of Helsinki.

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