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Circulating protein synthesis rates reveal skeletal muscle proteome dynamics

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Here, we have described and validated a strategy for monitoring skeletal muscle protein synthesis rates in rodents and humans over days or weeks from blood samples. We based this approach on label incorporation into proteins that are synthesized specifically in skeletal muscle and escape into the circulation. Heavy water labeling combined with sensitive tandem mass spectrometric analysis allowed integrated synthesis rates of proteins in muscle tissue across the proteome to be measured over several weeks. Fractional synthesis rate (FSR) of plasma creatine kinase M-type (CK-M) and carbonic anhydrase 3 (CA-3) in the blood, more than 90% of which is derived from skeletal muscle, correlated closely with FSR of CK-M, CA-3, and other proteins of various ontologies in skeletal muscle tissue in both rodents and humans. Protein synthesis rates across the muscle proteome generally changed in a coordinate manner in response to a sprint interval exercise training regimen in humans and to denervation or clenbuterol treatment in rodents. FSR of plasma CK-M and CA-3 revealed changes and interindividual differences in muscle tissue proteome dynamics. In human subjects, sprint interval training primarily stimulated synthesis of structural and glycolytic proteins. Together, our results indicate that this approach provides a virtual biopsy, sensitively revealing individualized changes in proteome-wide synthesis rates in skeletal muscle without a muscle biopsy. Accordingly, this approach has potential applications for the diagnosis, management, and treatment of muscle disorders.

Introduction
Disorders of muscle mass, quality, and function cause substantial and increasing morbidity and mortality. Sarcopenia, cachexia, and frailty are growing in importance in association with aging demographics worldwide (1, 2). Loss of skeletal muscle mass is a major target for drug development (3), but this field has been held back by the absence of simple translational biomarkers that can be used for diagnosis, prognosis, and monitoring. The synthesis and breakdown rates of skeletal muscle proteins of different classes are perturbed in muscle-wasting conditions (4) and increasing muscle protein synthesis is the primary metabolic mechanism of action of anabolic interventions proven to increase muscle mass and strength, such as resistance exercise and androgen treatment (5–10). Changes in mixed muscle protein synthesis rates occur very rapidly in response to anabolic interventions in humans (8, 11) — in advance of changes in muscle mass, strength, or performance (5, 9, 10) — and therefore would be ideal biomarkers for early assessment prediction and monitoring of treatment efficacy.

Translatable metrics of skeletal muscle protein turnover that can be applied routinely in therapeutic trials or in the clinic have, however, not been available. Blood- or urine-based tests of intracellular protein turnover in skeletal muscle would be particularly useful as biomarkers. An accurate, minimally invasive test of muscle protein dynamics might have applications for early detection of therapeutic response to therapeutic interventions, patient selection, and translating results from animal models to humans.

Fractional synthesis rates (FSRs) of mixed muscle proteins or protein subfractions are typically measured in rodents and humans through short-term infusions of stable isotope–labeled amino acids (12–16). These methods have demonstrated the anabolic effects of exercise (6, 7), dietary supplements (17, 18), or treatment with anabolic agents such as testosterone (8–10) or clenbuterol (19, 20). Although changes in FSR measurements precede longer-term responses of muscle mass and strength (5, 8–10), measurement of acute synthesis rates of mixed proteins in muscle has a number of fundamental limitations. First, the integrated effects on protein turnover of diet, activity, hormones, and medications over days or weeks are more relevant to muscle mass and function than turnover rates over hours, particularly for the long-lived structural and mitochondrial proteins that are characteristic of skeletal muscle. Second, broad interrogation of dynamics across proteins in different classes within the proteome is required to explore the coordinated control of expression and catabolism of different functional classes of proteins or to identify proteins involved in disease or interventions. And, third, measurement of muscle protein...
kinetics has previously required a tissue sample rather than being measurable noninvasively through a body fluid measurement.

A solution to the first problem is the use of oral intake of heavy water (\( ^2\text{H}_2\text{O} \)) in the outpatient setting to label newly synthesized proteins over periods of days, weeks, or months (21–28). We (25, 29–31) and others (32, 33) have shown that the second problem can be addressed by combining stable isotope label incorporation with tandem mass spectrometric–based proteomics techniques. In particular, isotope ratio measurements using liquid chromatography–mass spectrometry/mass spectrometry (LC-MS/MS) with quadrupole–time-of-flight (Q-ToF) instruments can be performed in the scan mode on trypsin-derived peptides with sufficient analytic accuracy to quantify synthesis rates of hundreds of proteins concurrently after relatively low-level in vivo \( ^2\text{H}_2\text{O} \) labeling (25, 29–31).

Here, to address the third problem noted above, we report the development and validation of blood test approach for measuring the integrated rate of muscle protein synthesis over days to weeks. Our goal was to develop a translatable mass spectrometric technique for monitoring skeletal muscle protein synthesis rates noninvasively, from a single blood sample that reflects the dynamics of the skeletal muscle proteome under basal and anabolic conditions. This minimally invasive method can be seen as a virtual biopsy for monitoring skeletal muscle protein synthesis rates. We asked whether the labeling kinetics of proteins in plasma that were synthesized in skeletal muscle, such as creatine kinase M-type (CK-M) and carbonic anhydrase 3 (CA-3) the dynamics of the skeletal muscle proteome, using \( ^2\text{H}_2\text{O} \) labeling combined with LC-MS/MS analysis (25, 29–33). In addition, the synthesis and turnover rates of multiple proteins across the proteome (i.e., proteome dynamics) had not previously been characterized in human skeletal muscle. We report that synthesis rates of proteins across the muscle proteome generally change in a coordinate manner in response to a sprint interval exercise training regimen in humans and to anabolic or catabolic interventions in rodents. Moreover, we report that labeling kinetics of plasma CK-M and CA-3 revealed these coordinate changes while detecting interindividual differences in muscle protein dynamics.

In combination with a technique for measuring skeletal muscle mass from a spot urine sample that we and our colleagues recently described (34–36), these translatable techniques have potential use as biomarkers in pharmacologic research and clinical management of disorders of skeletal muscle.

Results

Preclinical studies

Proteome dynamics in rat skeletal muscle after denervation and clenbuterol treatment. Proteome-wide synthesis rates were measured in rat gastrocnemius muscle by LC-MS/MS after 4 days of \( ^2\text{H}_2\text{O} \) administration to determine the broad effects of 2 interventions that are well established to alter skeletal muscle mass denervation and treatment with clenbuterol (20, 37). Sciatic nerve denervation produced a significant 30% decrease in weight of gastrocnemius muscle in vehicle-treated groups (contralateral: 623.5 ± 27.2 mg, denervation: 432.5 ± 37.6 mg, \( P < 0.001 \)), whereas clenbuterol treatment significantly increased the weight of muscle in denervated limb by 15% (vehicle: 432.5 ± 37.6 mg, clenbuterol: 498.5 ± 15.8 mg, \( P < 0.05 \)). Values represent mean ± SD of 4 rats per group and were significantly different by 2-way ANOVA.

FSRs met analytic criteria and were determined for each group (\( n = 4 \) per group) of animals, and there were 117 ± 23 proteins from the denervated limb, 114 ± 25 proteins from the contralateral limb, 122 ± 26 proteins from the vehicle-treated muscle, and 109 ± 20 proteins from the clenbuterol-treated muscle; the values represent the mean ± SD of number of protein FSRs measured in individual rat per group. For 75 of the proteins, synthesis rates were measurable in all 4 groups and therefore could be compared; these are represented as a heatmap (Figure 1A). Unsupervised hierarchical clustering analysis of protein FSRs spontaneously organized animals into 4 groups, which were identical to the 4 treatment groups. In addition, a principal component analysis (PCA) of the data revealed that muscle FSRs were sufficient to differentiate each of the groups from the others (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI79639DS1). Of special interest to our virtual biopsy approach, FSR of the cytosolic enzyme CK-M were significantly decreased by denervation and increased by clenbuterol treatment in both the contralateral and denervated limbs (Figure 1B). In contrast, FSR of another cytosolic enzyme, CA-3, was increased by clenbuterol treatment in contralateral limb and also increased by denervation (Figure 1C). FSRs of contractile myofibril proteins were significantly lower in the denervated limb compared with the contralateral limb, and treatment with clenbuterol significantly increased the FSRs of most proteins in both the denervated and control limbs (Figure 1D). A similar pattern was apparent for glycolytic enzymes, for which denervation uniformly decreased FSRs while clenbuterol increased FSRs (Figure 1E). Mitochondrial protein FSRs were significantly increased by clenbuterol only in the denervated limb but not in the contralateral limb, and they were not significantly altered by either clenbuterol or denervation alone (Figure 1F). Thus, the FSR signatures for the clenbuterol (anabolic) versus denervation (catabolic) groups were not entirely symmetric across all of these ontologies.

Virtual biopsy: FSR of CK-M and CA-3 in rat muscle and plasma after clenbuterol treatment. Next, we asked whether muscle protein FSRs could be measured in a minimally invasive manner in rats based on the FSR of a protein isolated from the bloodstream. CK-M is a cytosolic enzyme predominantly (>90%, ref. 38) present in skeletal muscle that can be detected in serum or plasma and is routinely released from damaged muscle, which has led to its use as a biomarker of rhabdomyolysis (39). CA-3 is virtually specific to skeletal muscle in humans and can also be detected in circulation (40, 41). In rats, CA-3 is an abundant cytosolic enzyme in both muscle and liver (42). We hypothesized that the FSRs of CK-M or CA-3, synthesized in skeletal muscle and isolated from plasma after in vivo labeling, might provide a virtual biopsy of the FSR of skeletal muscle proteins.

FSR of plasma and skeletal muscle CK-M and CA-3 were measured in rats at 3, 7, or 14 days of \( ^2\text{H}_2\text{O} \) labeling during clenbuterol treatment. At all time points, the fractional synthesis (\( f \)) of CK-M was higher in the gastrocnemius and quadriceps muscle
muscle samples (Supplemental Tables 1 and 2). Similar findings were observed for CA-3, with clenbuterol treatment producing a significant increase in f of CA-3 in both muscle and plasma compartments (Figure 3, A–C). Muscle tissue CA-3 values of f correlated highly significantly with f of other skeletal muscle proteins (Supplemental Tables 1 and 2) and with f of plasma CA-3 (Figure 3, D and E). The observations that both CK-M and CA-3 synthesis rates correlated well with synthesis rates of most proteins in muscle under anabolic conditions is consistent with the view that skeletal muscle tends to accrue structural and cytosolic proteins as an ensemble (43) and suggests that plasma CK-M and CA-3 synthesis rates should reflect the general synthesis rate of muscle structural and cytosolic proteins in the rat.

of clenbuterol-treated rats than in vehicle-treated animals (Figure 2, A and B). There was a highly significant correlation between f of muscle tissue CK-M and other muscle proteins measured in the tryptic digest of skeletal muscle samples (Supplemental Tables 1 and 2). The f of CK-M immunoprecipitated from plasma was also significantly increased after clenbuterol treatment (Figure 2C). Most importantly, there was a highly significant correlation (r = −0.98) between the f of CK-M isolated from plasma and the f of CK-M measured in gastrocnemius (Figure 2D) and quadriceps (Figure 2E) muscle tissue from the same animals, consistent with the principle that >90% of plasma CK-M is derived from skeletal muscle (38). Plasma CK-M f also correlated highly significantly with the f of other proteins measured in the muscle samples (Supplemental Tables 1 and 2). Similar findings were observed for CA-3, with clenbuterol treatment producing a significant increase in f of CA-3 in both muscle and plasma compartments (Figure 3, A–C). Muscle tissue CA-3 values of f correlated highly significantly with f of other skeletal muscle proteins (Supplemental Tables 1 and 2) and with f of plasma CA-3 (Figure 3, D and E). The observations that both CK-M and CA-3 synthesis rates correlated well with synthesis rates of most proteins in muscle under anabolic conditions is consistent with the view that skeletal muscle tends to accrue structural and cytosolic proteins as an ensemble (43) and suggests that plasma CK-M and CA-3 synthesis rates should reflect the general synthesis rate of muscle structural and cytosolic proteins in the rat.

Figure 1. Proteome dynamics in rat gastrocnemius muscle. (A) Heatmap of FSRs (% day⁻¹) of 75 proteins in rat muscle measured in n = 3 rats per group after denervation and clenbuterol treatment, with each horizontal line representing z-scaled FSR of an individual protein. (B) Effect of denervation surgery and clenbuterol treatment on FSR of CK-M, mean ± SD, n = 3–4/group. *P < 0.05 'Den-Veh' vs. 'Con-Veh', †P < 0.05 'Den-Clen' vs. 'Den-Veh', 2-way ANOVA. (C) Effect of denervation surgery and clenbuterol treatment on FSR of CA-3, mean ± SD, n = 3–4/group. *P < 0.05 'Den-Veh' vs. 'Con-Veh', †P < 0.05 'Con-Clen' vs. 'Con-Veh', 2-way ANOVA. (D–F) Effect of denervation surgery and clenbuterol treatment on the FSRs of myofibril, glycolytic, and mitochondrial proteins. Names are listed in the order they appear on the scatter plots for each class of proteins. Data represent mean ± SD, n = 3–4/group, * corrected P < 0.05 'Den-Veh' vs. 'Con-Veh', † corrected P < 0.05 'Con-Clen' vs. 'Con-Veh', 2-way ANOVA with Holm-Sidak correction. Con, control; Veh, vehicle; Clen, clenbuterol; Den, denervation.
accessed programmatically from the Database for Annotation, Visualization and Integrated Discovery (DAVID) v6.7 (44, 45), a paired \( t \) test of protein FSRs by gene ontology was performed. A total of 8 nonredundant biological processes were enriched with significant differences (\( P < 0.05 \) in paired \( t \) tests for proteins, with Benjamini-Hochberg multiple test corrections) in mean protein FSR when comparing kinetics of the muscle proteome from subjects who were sedentary verses those who underwent sprint exercise. Proteins assigned to several gene ontologies were significantly affected following sprint exercise, including the 29 proteins classified as being involved in glucose metabolic processes, the 17 proteins involved in regulation of apoptosis, the 16 proteins involved in striated muscle contraction, and the 12 proteins involved in cellular respiration (Figure 4B); individual protein data are shown in Supplemental Table 3B. We also performed a between-group unpaired \( t \) test on 67 proteins that were measured in all subjects. Following Benjamini-Hochberg multiple test corrections, there were 20 proteins that were significantly (FDR < 0.2) different between the groups (Figure 4C), reflecting the variability of protein metabolic response to exercise. The molecular kinetic findings of higher FSR changes in glycolytic enzymes and contractile proteins, compared with smaller changes in mitochondrial proteins, may provide insight regarding the functional consequences of this widely used sprint exercise program.

**Virtual biopsy: CK-M and CA-3 FSRs in plasma compared with muscle in humans.** Our primary goal in these studies was to develop proteome-wide synthesis rates were measured in quadriceps muscle biopsies from 11 healthy human subjects, 6 of whom had gone through a sprint-interval exercise program as described elsewhere (25), while the other 5 subjects were sedentary controls. Subjects drank 50 ml of \( ^{2} \text{H}_{2} \text{O} \) (70% enriched) twice daily for 3 weeks after an initial 1-week priming protocol of 50 ml 3 times a day. FSRs were measured for 515 proteins, of which FSRs for 273 proteins were measurable in \( \geq 2 \) subjects per group and could therefore be compared. FSRs ranged from a minimum of 2.2% per week to a maximum of 75% per week (Supplemental Table 3A). Sprint exercise produced a substantial increase in FSRs of most proteins analyzed. The FSRs of 139 individual proteins that were measured at least in 60% of the subjects in each group are shown in a heatmap (Figure 4A). Interindividual differences in FSR are apparent and were consistent across most protein through each subject’s sampled proteome, reflecting a characteristic, individualized response to exercise. Pearson cross correlation coefficient analyses revealed that FSRs in muscle were more similar across proteins measured in sedentary subjects (average \( r = 0.883 \)) compared with subjects that participated in the sprint exercise program (average \( r = 0.722 \)).

The mean FSRs of proteins in sedentary and sprint groups were further compared at the gene ontological level. Using a data analysis program to group proteins based on their ontology terms

**Human studies**

**Proteome dynamics in human skeletal muscle: effects of sprint exercise.** Proteome-wide synthesis rates were measured in quadriceps muscle biopsies from 11 healthy human subjects, 6 of whom had gone through a sprint-interval exercise program as described elsewhere (25), while the other 5 subjects were sedentary controls. Subjects drank 50 ml of \( ^{2} \text{H}_{2} \text{O} \) (70% enriched) twice daily for 3 weeks after an initial 1-week priming protocol of 50 ml 3 times a day. FSRs were measured for 515 proteins, of which FSRs for 273 proteins were measurable in \( \geq 2 \) subjects per group and could therefore be compared. FSRs ranged from a minimum of 2.2% per week to a maximum of 75% per week (Supplemental Table 3A). Sprint exercise produced a substantial increase in FSRs of most proteins analyzed. The FSRs of 139 individual proteins that were measured at least in 60% of the subjects in each group are shown in a heatmap (Figure 4A). Interindividual differences in FSR are apparent and were consistent across most protein through each subject’s sampled proteome, reflecting a characteristic, individualized response to exercise. Pearson cross correlation coefficient analyses revealed that FSRs in muscle were more similar across proteins measured in sedentary subjects (average \( r = 0.883 \)) compared with subjects that participated in the sprint exercise program (average \( r = 0.722 \)).

The mean FSRs of proteins in sedentary and sprint groups were further compared at the gene ontological level. Using a data analysis program to group proteins based on their ontology terms

**Figure 2. Effects of clenbuterol treatment on f of CK-M in rat muscles and plasma.** (A) CK-M f (% in gastrocnemius muscle of rats treated with clenbuterol for 3, 7, or 14 days (mean ± SD, \( n = 4 \)/group, \( \times P < 0.05 \) clenbuterol vs. vehicle, 2-way ANOVA). (B) CK-M f in quadriceps muscle of rats treated with clenbuterol for 3, 7, or 14 days (mean ± SD, \( n = 4 \)/group, \( \times P < 0.05 \) clenbuterol vs. vehicle, 2-way ANOVA). (C) CK-M f in plasma of rats treated with clenbuterol for 3, 7, or 14 days (mean ± SD, \( n = 4 \)/group, \( \times P < 0.05 \) clenbuterol vs. vehicle, 2-way ANOVA). (D) Significant correlation (Pearson correlation \( r > 0.95, P < 0.001 \)) between fractional syntheses of plasma CK-M compared with that of gastrocnemius CK-M. (E) Significant correlation (Pearson correlation \( r > 0.95, P < 0.001 \)) between fractional syntheses of plasma CK-M compared with that of quadriceps CK-M.
In order to determine whether plasma CK-M or CA-3 FSRs can be used as virtual biopsies of muscle CK-M, CA-3, and general protein synthesis rates in humans, FSR of CK-M and CA-3 were compared in plasma and a muscle biopsy specimen (vastus lateralis) for the 17 healthy subjects. All subjects drank H₂O₂ for 4 weeks. CK-M FSR in human muscle ranged from 3.9%–11.7% per week in young male and female sedentary and exercising subjects compared with FSR in plasma of the same subjects of 4.5%–12.3% per week. There was a highly significant correlation between the FSR values of CK-M isolated from muscle and plasma ($r > 0.89$, Figure 5B), similar to the observation in preclinical studies. Similarly, FSR of CA-3 in human muscle ranged from 2.8%–9.1% per week, compared with FSR in plasma of 4.4%–9.6% per week, with a very highly significant correlation ($r > 0.87$, Figure 5C).

Correlation of CK-M and CA-3 FSRs with FSRs of human muscle proteins. We also determined the correlations between FSRs of CK-M and CA-3 in the 17 subjects correlated closely and highly significantly ($r = 0.84–0.94$, $P < 0.0001$) with that of gastrocnemius CA-3. (E) Significant correlation (Pearson correlation $r > 0.88$, $P < 0.001$) between fractional syntheses of plasma CA-3 compared with that of quadriceps CA-3.

To establish the time course of a muscle-derived protein in plasma, we first measured the FSR of CK-M isolated from human plasma samples in 3 male subjects who ingested H₂O₂ daily for 1–7 weeks. CK-M was immunoprecipitated from plasma using an anti–CK-M specific antibody, followed by gel fractionation, in-gel trypsin digestion, and LC-MS/MS analysis. The F of CK-M measured at each time point was fit to a monoeXponential rise to plateau equation ($F = 1-e^{-kt}$). The rate constant, $k$, (synonymous with FSR) was 1.0% ± 0.2% per day (Figure 5A). The curve fits were consistent with a plateau value close to 100%, and any single time point collected between 1–7 weeks was sufficient to reveal the FSR value calculated by the curve fit (mean FSR from individual time points = 1.0% ± 0.1% per day).
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Technical advance

7, A and B, and Supplemental Table 4). Since 2 of the subjects in the sprint group had high FSRs, correlations were also performed excluding these subjects, and these remained significant ($r = -0.6$–$0.8$, $P < 0.05$) for most proteins. These results indicate that in humans, as observed in rats, plasma CK-M and CA-3 synthesis rates reflect global muscle protein synthesis rates and detect the coordinated effects of an anabolic intervention on muscle protein dynamics.

Discussion

There were 2 primary goals of this work. The first was to develop a minimally invasive method for measuring integrated rates of skeletal muscle protein synthesis over a period of days to weeks based on a single blood measurement — i.e., without the need for a physical sample of the tissue. Muscle biopsies are not likely to be widely used in the clinical setting, so a blood test of muscle protein synthesis rates that could be monitored over time would be attractive.
for drug development, as well as clinical management of disorders of muscle, including sarcopenia and cachexia. Because changes in muscle protein synthesis rates have been shown to occur rapidly in response to anabolic therapies and to predict subsequent changes in muscle mass, strength, and performance (5, 8–10), the availability of a blood test for skeletal muscle protein synthesis rates would greatly simplify translational drug development and monitoring of treatment efficacy in this growing, unmet medical need.

We describe here an approach that we call a virtual biopsy, based on a simple concept: if proteins can be identified in a body fluid that derive essentially exclusively from a single tissue, and if the rate-determining or slow kinetic step for such proteins is in the tissue (i.e., the half-life in the body fluid is shorter than in the tissue of origin), then the measured FSR of the protein isolated from the body fluid will reflect the FSR of the protein in the tissue of origin.

Here, we chose CK-M and CA-3 as candidate proteins to represent skeletal muscle protein turnover. CK-M and CA-3 were attractive candidates in principle because these circulating proteins are known to be >90% derived from skeletal muscle in humans (38, 40, 41), and their clearance from plasma is relatively fast in healthy subjects (46) while their half-lives in muscle are slow (~2 months, as shown in Figure 4). These assumptions needed to be validated experimentally in rodents and humans. We show that FSR of muscle CK-M and plasma CK-M are essentially identical, and this is also true for FSR of muscle CA-3 and plasma CA-3, meeting the central principle of a virtual biopsy. Perhaps more importantly, FSRs of plasma and muscle CK-M and CA-3 correlated highly with FSRs of a large number of muscle proteins across multiple gene ontologies. The latter finding is consistent with the concept that muscle cytosolic and structural proteins tend to be synthesized as an ensemble when muscle mass increases or diminishes; i.e., the proportional content of muscle cells does not change when mass changes under anabolic and catabolic conditions (43). Repeated measurements of CK-M synthesis on blood samples at different time points gave reproducible and consistent results in subjects at steady-state for muscle mass, suggesting that monitoring of muscle FSR over time should be feasible to track the early effect of interventions directed at increasing muscle mass. The coordinated assembly of muscle proteins (43) and the confluence of muscle protein synthesis rates further supports the idea of using a representative and accessible protein, such as CK-M or CA-3 in plasma, to represent general protein dynamics in muscle.

It should be noted that CK-M and CA-3 are only 2 possible candidates for muscle-derived proteins that can be isolated from plasma to measure tissue protein synthesis rates, and they are particularly useful to track effects of anabolic interventions. In as much as no single protein, whether in the cell or in the circulation, can serve as the sole read-out for changes in all classes of muscle protein turnover, these cytosolic proteins may not accurately reflect all the protein kinetic changes in other compartments, such as mitochondria, especially when there are differential subcellular effects of interventions. There may be several other blood proteins that are similar or superior for this virtual biopsy purpose, making it feasible to identify a panel of blood proteins for monitoring different aspects of muscle protein dynamics. The finding that 2 different proteins in plasma that are believed to derive primarily from skeletal muscle in fact showed very similar FSRs in plasma and muscle strongly supports the general principle of virtual biopsy and suggests that other proteins will likely be identified. Interestingly, it may be possible to measure FSRs of muscle-specific mitochondrial proteins in the

![Figure 5. CK-M and CA-3 synthesis in human plasma and skeletal muscle. (A) The f (%) of CK-M in 4 subjects after continuous $^2$H$_2$O labeling for several weeks. (B) Significant correlation (Pearson correlation $r = 0.89$, $P < 0.0001$, $n = 17$) between FSR (% week$^{-1}$) of CK-M measured in the plasma and in the skeletal muscle of the same subject. (C) Significant correlation (Pearson correlation $r = 0.88$, $P < 0.0001$, $n = 17$) between FSR of CA-3 measured in the plasma and in the skeletal muscle of the same subject.]
Figure 6. Muscle CK-M and CA-3 synthesis correlate with skeletal muscle protein synthesis. (A) Significant correlation (Pearson correlation $r > 0.8$, $P < 0.0001$, $n = 16-17$) between FSR (% week$^{-1}$) of CK-M and FSRs of myofibrillar proteins, both measured in skeletal muscle of the same subject. (B) Significant correlation (Pearson correlation $r > 0.8$, $P < 0.0001$, $n = 16-17$) between FSR of CA-3 measured in plasma and FSRs of myofibrillar proteins measured in skeletal muscle of the same subject.
circulation: 2 peptides from COX8B have been identified only in skeletal and cardiac muscle of mice (MitoCarta1.0 database, ref. 47). Although this protein has not been identified in humans, 2 peptides from COX6A2, which is present in humans, were also identified as muscle specific among the 14 mouse tissues studied (MitoCarta2.0 database, ref. 48). We have recently described a similar virtual biopsy approach for measuring hepatic fibrogenesis rates from blood samples by measuring FSRs of extracellular matrix-related proteins (such as lumican or TGFβ-induced protein) that escape from liver into the circulation (49). We believe that the underlying concept presented here may have broad applications, building on the great sensitivity of tandem mass spectrometry when combined with stable isotope labeling.

It is important to point out the differences between measuring concentrations versus synthesis rates of plasma proteins derived from muscle. Conceptually, the level of a protein in plasma has no fundamental relationship to its synthesis rate back in the tissue of origin but is a function of factors, such as release rate from the tissue and clearance rate from the blood. Because mass spectrometric analysis compares an internal ratio among different masses within a molecule, yield or recovery of the molecule do not, in principle, affect kinetic calculations and are therefore not subject to influence of tissue leakage or clearance from plasma. This feature provides a real advantage of kinetic measurements for blood-based assays.

Our second objective was to measure the synthetic rates of large numbers of proteins reproducibly across a variety of gene ontologies in muscle tissue from humans and experimental animals, in response to anabolic or catabolic interventions. This goal required an analytic approach that goes beyond small numbers of targeted proteins measured one at a time by LC-MS/MS (50–52). We have found that Q-ToF MS/MS instruments achieve the highest analytic accuracy and precision for quantifying mass isotope ratios in hundreds or thousands of peptides concurrently, which allows accurate and reproducible protein kinetic measurements from a single time point (25, 29–31). Using this approach, we have identified characteristic proteome dynamic signatures of several conditions, including calorie restriction in liver (31), experimental fibrosis in lung (29), proliferating neural stem cells in vitro (53), neuroinflammation in mouse brain and cerebrospinal fluid (54), and FSRs of human plasma proteins (30), but the detailed proteome dynamic response in human muscle to anabolic or catabolic conditions had not previously been measured.

We demonstrate here analytically robust kinetic measurements, as evidenced by the consistent changes within subjects and within functionally related classes of proteins in response to interventions. The labeling protocol is relatively nonburdensome, involving oral intake of 2H2O in drinking water for 4–7 days in rodents and 2–3 daily doses for 1–4 weeks in humans. We report on the turnover rates of approximately 500 proteins in human skeletal muscle and approximately 100 proteins in rat skeletal muscle. The former number is larger in part because of differences in protein identification due to a larger Swiss-Prot human proteome database and also because we used a more sensitive model of the instrument (Q-ToF 6550) for the human analyses.

Technically, stringent analytic accuracy for isotope ratio measurements is required to quantify relatively low levels of deuterium incorporation during 2H2O administration in humans and preclinical models (25, 29–31). Because the stringent analytic filtering criteria we have found are required for reliable kinetic measurements (30), the proteome-wide kinetic measurements described here using Q-ToF instruments quantify fewer proteins than can be identified with static proteomic measurements of protein content, especially when highly sensitive Orbitrap instruments are used for the latter (55). This is because a large fraction of identified peptides do not meet analytic criteria for quantitative accuracy of isotope ratio measurements. In addition, Orbitrap instruments to date have not been able to achieve sufficiently accurate isotope ratio measurements on peptides in the scan mode (56) to allow usable proteome-wide kinetic measurements at the low level deuterium enrichments that are achieved in human studies, which is why we have developed our current proteome dynamic approach using Q-ToF instruments. Having kinetic values for up to several hundred proteins in a tissue spans a wide variety of gene ontologies and functions, however (Figures 1 and 3 and refs. 29, 31) and allows broad conclusions about cellular response.

Earlier studies have documented a reduction in mixed muscle protein synthesis rates and content in skeletal muscle tissue (20, 57) or in mixed myofibril protein subfractions (37), after denervation, using radioactive amino acid tracers. More recent proteomic approaches have quantified changes in the abundances of specific proteins (58–60). Our quantitative profile of FSR changes of >100 specific proteins in response to denervation extend these findings and are consistent with gene expression changes in denervated muscle reported for structural proteins, enzymes involved in glycolysis, and mitochondrial ATP synthesis (61). Interestingly, the protein anabolic dynamic signature for clenbuterol treatment was not in all respects reciprocal to the signature for denervation. Clenbuterol produced a strong response of increased FSRs for cytosolic and structural proteins, as expected (20, 57), but its effects on mitochondrial proteins were limited to only the denervated limb, suggesting differential mechanisms and a stronger anabolic response under conditions of muscle atrophy. In addition, consistent with previous reports that denervation induces CA-3 in rat skeletal muscle (62, 63), we also observed that denervation increased CA-3 synthesis, whereas most cytosolic protein FSRs were reduced. Accordingly, in denervated rats, muscle or plasma CA-3 synthesis rates are not representative of reduced synthesis rates of most muscle cytosolic proteins.

In humans, the proteome dynamics revealed somewhat unexpected results for the skeletal muscle proteostatic response to sprint interval training (SIT), a program that involves repetitive 30-second bouts of intensive exercise (64, 65). While several reports have documented increased glycolytic and oxidative enzyme activities in response to SIT (64–67), our results indicate that higher synthesis rates were observed for glycolytic enzymes and cytosolic proteins but that mitochondrial proteins in general did not exhibit higher FSRs in subjects on sprint compared with sedentary regimens. These protein dynamic findings provide molecular kinetic insight into the physiologic consequences of this exercise training regimen, suggesting a primary functional impact on the synthesis of proteins related to muscle anaerobic power and contractile strength. Proteins involved in regulation of apoptosis and oxidative stress, including heat-
Figure 7. Plasma CK-M and CA-3 synthesis rates provide a virtual biopsy of skeletal muscle protein synthesis. (A) Significant correlation (Pearson correlation \( r > 0.7, P < 0.0005, n = 16-17 \)) between FSR (% week\(^{-1}\)) of CK-M measured in the plasma and FSRs of myofibrillar proteins measured in skeletal muscle of the same subject. (B) Significant correlation (Pearson correlation \( r > 0.7, P < 0.0005, n = 16-17 \)) between FSR of CA-3 measured in plasma and FSRs of myofibrillar proteins measured in skeletal muscle of the same subject.
The availability of a blood test for monitoring muscle protein FSR and treatment efficacy in drug development or patient management, have a number of possible uses in drug development and for advancing our understanding of nutritional and exercise effects on muscle biology, as well as potentially for medical diagnosis, prognosis, and monitoring.

Methods

**Animals.** Male Sprague-Dawley (SD) rats (250–300 g; Charles River Laboratories), were used for these studies. Animals were housed in a climate-controlled environment with a 12-hour light/dark cycle and were fed standard rodent chow and water ad libitum.

**Surgery.** Sciatic denervation surgery was used as a model for muscle atrophy and wasting as previously described (74). Animals were placed under isoflurane anesthesia, and buprenorphine was administered at the time of surgery. The sciatic nerve of one leg was exposed in the thigh region by blunt forceps reflection of the muscle. One hind limb was denervated with a hemostat by crushing the sciatic nerve immediately proximal to the division of the peroneal and tibial branches for 30 seconds. The skin was closed with wound clips. Denervation was known to be complete by observing postoperative disuse of the lower limb and loss of a withdrawal response to a pinch in the sciatic distribution area. The contralateral leg was sham operated, and its muscles were used as internal controls for the denervated tissues.

**Drug treatment.** Animals (n = 4 per group) started drug treatment by oral gavage 3 days after denervation surgery with either vehicle (water) or clenbuterol HCL (4 mg/kg) dosed once daily for 4 days. In a separate time-course experiment, rats were gavaged daily with vehicle (water) or clenbuterol HCL (4 mg/kg) for either 3, 7, or 14 days.

**2H2O labeling protocol in rats.** Animals were labeled with 2H2O concurrent with vehicle or clenbuterol treatment, which was 4 days in the denervation study and either 3, 7, or 14 days in the time-course study. Animals received a priming i.p. bolus of 35 ml/kg 0.9% NaCl in 99.9% 2H2O and were maintained on 8% 2H2O in drinking water until sacrifice. This regimen results in steady-state values of about 5% excess 2H enrichment of body water.

**Tissue harvesting.** Animals were euthanized by CO2 asphyxiation. Blood was drawn by cardiac puncture and centrifuged at 2,000 g for 10 minutes to obtain plasma, which was stored at −20°C until processing. The gastrocnemius and quadriceps muscles were dissected out of each leg and stored at −20°C until further processing.

**2H2O labeling protocol in humans.** Deuterium labeling of newly synthesized proteins was achieved by oral consumption of 2H2O (70%; Cambridge Isotope Laboratories) for 4 weeks using protocols described previously (24, 25, 30). A target of 1%–2% enrichment was achieved during a 1-week priming stage, when subjects consumed 50 ml of 70% 2H2O three times a day for a total of 150 ml/day, which was maintained for 3 weeks with a dose of 50 ml of 70% 2H2O two times a day for a total of 100 ml/day. Subjects in the time-course study consumed 50 ml of 70% 2H2O three times a day for a total of 150 ml/day,
which was maintained for a maximum of 7 weeks with a dose of 40 ml of 70% \( ^{2}H_{2}O \) two times a day for a total of 80 ml/day.

**SIT.** Participants completed 9 sessions of SIT over 3 weeks. Each session consisted of four to eight 30-second bouts of maximal efforts on a stationary cycle ergometer (Velotron, RacerMate Inc.) against a resistance equal to 7.5% of participant body mass separated by 4 minutes of active recovery; to facilitate recuperation, each session was separated by 1-2 days. The first and ninth sessions both consisted of four 30-second bouts to allow for comparisons to be made between these sessions.

**Sample collection.** Saliva swabs were collected periodically during oral \( ^{2}H_{2}O \) consumption and stored at ~80°C until analysis. Venous blood was collected either during weekly blood draws or at the end of SIT training, and plasma was separated by centrifugation and stored at ~80°C. While the subjects were under local anesthesia (1% lidocaine), muscle biopsy samples (~100-150 mg) of the vastus lateralis were removed using a 5-mm Bergström needle with manual suction and then immediately frozen in liquid nitrogen and stored at ~80°C.

**Body water enrichment analysis.** Body water enrichment was determined from saliva swabs or plasma samples in humans and in rat plasma samples. Aliquots of plasma or saliva were diluted 1:200 and placed into the caps of inverted sealed screw-capped vials for overnight distillation at 80°C. Body water \( ^{2}H_{2}O \) enrichments were determined by direct measurement of deuterium mole percent excess (MPE) in water distilled from the blood plasma. MPE was measured against a \( ^{2}H_{2}O \) standard curve using laser water isotope analyzer (LGR).

**SDS-PAGE fractionation, Coomassie blue staining, and in-gel trypsin digestion of muscle proteins.** Human or rat muscle tissues were prepared for LC/MS analysis using methods described previously (31). Briefly, 10-30 mg muscle samples were homogenized in M-PER reagent (Thermo Scientific) at 100 mg/ml with 1× protease inhibitor cocktail (Thermo Scientific). Protein (250 μg) from prepared homogenates was uniformly reduced via incubation in 5 mM Tris(2-carboxyethyl)phosphine (TCEP) and SDS-PAGE sample loading buffer for 5 minutes at 95°C, alkylated via incubation in 15 mM iodoacetamide for 1 hour at room temperature, and fractionated by SDS-PAGE (Invitrogen). The gel bands corresponding to 10 discrete molecular weight regions were excised from Coomassie blue-stained gels and digested overnight with trypsin (Proteomics Grade, Sigma-Aldrich) at 37°C. The peptides were extracted from the gel, dried, and reconstituted in 3% acetonitrile/0.1% formic acid for LC/MS analysis.

**Immunoprecipitation of CK-M and CA-3 from plasma, SDS-PAGE fractionation, and in-gel trypsin digestion.** CK-M was immunoprecipitated from 500 μl-2 ml rat or human plasma using 20 μg of goat anti–CK-M polyclonal antibody (CalBioreagents, P195) conjugated to 1 mg epoxy Dynabeads (Invitrogen). Samples were incubated for 60 minutes at room temperature (RT), and the bound CK-M was eluted in sample buffer and prepared for SDS-PAGE fractionation and Coomassie blue staining as described above. The gel bands corresponding to 37-50 kD were excised and subjected to overnight trypsin (Proteomics Grade, Sigma-Aldrich) digestion at 37°C. The peptides were extracted from the gel, dried, and reconstituted in 3% acetonitrile/0.1% formic acid for LC/MS analysis. CA-3 was immunoprecipitated from 2 ml rat or human plasma using 20 μg of goat anti–CA-3 polyclonal antibody (R&D Systems, AF2185) conjugated to 1 mg epoxy Dynabeads (Invitrogen). Samples were incubated for 60 minutes at RT, and the bound CA-3 was eluted in 30% acetonitrile, 0.5% formic acid (pH ~2.5), followed by in-solution trypsin digestion for LC/MS analysis.

**LC/MS analysis.** Trypsin-digested peptides were analyzed on a 6520 or 6550 Q-ToF mass spectrometer with 1260 Chip Cube ESI source (Agilent Technologies). Peptides were separated using a Polaris HR chip (Agilent Technologies, G4240-62030) consisting of enrichment and analytical columns packed with Polaris C18-A stationary phase. Mobile phases were 5% v/v acetonitrile and 0.1% formic acid in water and 95% acetonitrile and 0.1% formic acid in water. Peptides were eluted at 350 nL/minute flow rate with an 18 minutes LC gradient (for IP samples) or a 27 minutes LC gradient (for gel band samples). Each sample was analyzed once for protein/peptide identification in data-dependent MS/MS mode and once for peptide isotope analysis in MS mode. Acquired MS/MS spectra were extracted and searched using Spectrum Mill Proteomics Workbench software (version B.04.00, Agilent Technologies) and either a rat or human protein database (http://www.uniprot.org/, release 2013_05). Search results were validated with a global false discovery rate of 1%. A filtered list of peptides was collapsed into a nonredundant peptide formula database containing peptide elemental composition, mass, and retention time. This was used to extract mass isotope abundances (MO–M3) of each peptide from MS-only acquisition files with Mass Hunter Qualitative Analysis software (version B.05.00, Agilent Technologies). Software was developed at KineMed Inc. to calculate peptide elemental composition and curve-fit parameters for predicting peptide isotope enrichment (EM0) based on precursor body water enrichment (p) and the number (n) of amino acid C–H positions per peptide actively incorporating hydrogen (H) and deuterium (D) from body water. Subsequent data handling was performed using Microsoft Excel templates, with input of precursor body water enrichment for each subject, to yield FSR data at the protein level. FSR data were filtered to exclude protein measurements with fewer than 2 peptide isotope measurements per protein. Details of FSR calculations and data filtering criteria were described previously (30, 31).

**Statistics.** Two-way ANOVA with post-hoc Holm-Sidak correction for multiple comparisons (GraphPad Prism) were performed to determine the differences between the groups for the rat denervation and clenbuterol study protein FSRs, as well as CK-M and CA-3 FSRs in muscle and plasma from the clenbuterol time-course study. Differences were considered significant at corrected \( P < 0.05 \). Pearson correlation analysis (GraphPad Prism) was performed to correlate the f of CK-M and CA-3 in the muscle to that measured in the muscle. In addition, correlation analysis was also performed with f of plasma CK-M and CA-3 with that of several myofibril and cytosolic proteins measured in the muscle. Correlations were considered significant at \( P < 0.05 \). Heatmaps, hierarchical clustering, PCA analysis, and Pearson cross-correlation analysis of muscle protein FSR data were performed using Inferno\(^{10}\) for Proteomics. An in-house data analysis tool was developed to query and statistically analyze ontology terms of human muscle proteins, accessed programmatically from the DAVID v6.7 Gene ontology terms were retrieved based on an initial search of the proteins identified in the experimental datasets. The resulting terms were organized by categories (biological process, cellular component, and molecular function) and by levels (1-5). Duplicate terms within categories occurring at multiple levels were filtered to include only unique ontology terms at the lowest level. The median, standard deviation, and number of matching proteins were calculated for each
ontology term, based on the corresponding proteins from experimental data. Paired 2-tailed t tests with Benjamini-Hochberg multiple test corrections were then used to determine which terms were significantly enriched between experimental groups (corrected \( p < 0.05 \)). Significant terms were then further filtered by calculating the intersection of matching proteins within each ontology term in the same category and level, removing those terms with a minimum 80% intersection with other term(s) and retaining terms with the highest number of proteins. In addition, a between-group, unpaired 2-tailed t test was also performed on 67 human muscle protein FSRs measured in all subjects; differences were considered significant with a false discovery rate of 0.2 after a Benjamini-Hochberg procedure was performed to adjust for the multiple comparisons.

Study approval. All animal studies were carried out according to NIH guidelines for the care and use of laboratory animals and received prior approval by the KineMed Animal Care and Use Committee. All human procedures and protocols described in the clinical studies were approved by the Institutional Review Boards at either Colorado State University (17 subjects: 7 males, 10 females) or BioMed IRB (San Diego, California, USA) (4 males). Each volunteer was informed of the potential risks and benefits and provided written consent before participating. The study followed the guidelines set forth by the Declaration of Helsinki.

Author contributions
MS, SMT, CB, KLH, BFM, and MKH designed the studies. CLK performed studies and sample analyses. MS, TEA, WEH, KWL, MC, and JCP developed methods. MS, TEA, WEH, KWL, MC, JCP, and MKH analyzed data. MS, CLK, WEH, TEA, and MKH wrote the manuscript.

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