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**A MURINE MODEL OF GLUCOCORTICOID MYOPATHY**

**ALLEVIATION USING ANDROGEN THERAPY**

by

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*”Se questa non piace, non voglio più scrivere di musica.”*

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**A MURINE MODEL OF GLUCOCORTICOID MYOPATHY**

**ALLEVIATION USING ANDROGEN THERAPY**

# NICOLAE LUCIAN SANDOR

Boston University School/Division of Graduate Medical Sciences, 2015

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# Abstract

Glucocorticoids (GC) are used widely for the treatment of a large number of inflammatory conditions. Loss of muscle mass and muscle weakness are common complications of GC therapy. Androgen therapy has been suggested to reverse

GC-associated muscle loss (GAML), but evidence of its effectiveness is lacking.

Here, I established a mouse model of GAML. Young adult male mice receiving 0.25 mg/day dexamethasone (D) s.c. daily, for a week, lost 3% of their body weight. Through NMR lean body mass quantification and muscle dissection, a loss of more than 10% of their muscle mass was lost. More than half of the muscle loss was reversed by co-administration of 0.7 mg/day testosterone (T). This is the first mouse model of GAML alleviation by T.

Intramuscular atrogene expression and proteasome catalytic activity were upregulated by D and suppressed by T co-administration. T co-administration caused intramuscular downregulation of atrogene-activating Foxo transcription factors. Intramuscular pro-autophagic REDD1 and Klf15 were repressed by T co-administration. T co-administration reduced the autophagosome-characteristic lipidated form of LC3B. Translation regulators 4E-BP, eIF3f, and eIF2 did not change significantly as a result of androgen co-administration. Calpains activity and levels were unchanged by D and T.

C2C12 differentiated myotubes were used to determine the effects of T and D on protein synthesis and degradation. Myotube diameters were reduced by D, while T co-administration suppressed D effect. Protein degradation was increased by 24-hour D treatment. D-stimulated protein degradation was inhibited by proteasome inhibitor MG132, and, to a lesser degree, by lysosome inhibitor chloroquine. T co-administration returned protein degradation to basal levels. Protein synthesis response to D and T did not correlate with the observed phenotypes.

In vivo, D reduced intramuscular IGF-I expression, an effect reversed by T co-administration. In C2C12, inhibition of IGF-1R signaling with picropodophyllin did not modify T effects.

In conclusion, T protective action in GAML is mainly anti-catabolic, through reversal of proteasome and autophagosome upregulation induced by D. T stimulates a potentially protective intramuscular IGF-I response. Different models are needed to determine the role of protein synthesis and of IGF-I in

GAML.

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# Abbreviations

|  |  |
| --- | --- |
| 24HUC | 24-hour urine cortisol |
| 3MH | 3-methylhistidine |
| 4E-BP | eIF4E binding protein |
| AAP | adrenal androgen precursor |
| AAS | anabolic androgenic steroids |
| ACTH | adrenocorticotropic hormone; |
| ActRIIB | activin A receptor, type IIB |
| AMPK | adenosine monophosphate kinase |
| ANOVA | analysis of variance |
| AP-1 | activator protein 1 |
| AR | androgen receptor |
| ATF4 | activating transcription factor 4 |
| Atg12 | autophagy-related 12 |
| BCA | bicinchoninic acid |

|  |  |
| --- | --- |
| BCAA | branched-chain amino acids |
| Bcl3 | B-cell leukemia/lymphoma 3 |
| bFGF | basic fibroblast growth factor |
| BSA | bovine serum albumine |
| C/EBP | CCAAT-enhancer-binding protein |
| Comb | combination of Dexa and Testo |
| COPD | chronic obstructive pulmonary disease |
| CS | Cushing’s syndrome |
| CSA | cross-sectional area |
| CT | computed tomography |
| Ct | cycle threshold |
| D | dexamethasone-only group |
| Dexa | dexamethasone |
| DHEA | dehydroepiandrosterone |
| DHEAS | dehydroepiandrosterone sulfate |
| DHT | dihydrotestosterone |
| DMEM | Dulbecco’s Modified Eagle Medium |
| DMSO | dimethyl sulfoxide |
| DT | dexamethasone and testosterone group |

|  |  |
| --- | --- |
| DTT | dithiothreitol |
| EC50 | half maximal effective concentration |
| EDTA | ethylenediaminetetraacetate |
| eIF2 | eukaryotic initiation factor 2 |
| eIF2B | eukaryotic initiation factor 2B |
| eIF3-f | eukaryotic initiation factor 3f |
| eIF4E | eukaryotic translation initiation factor 4E |
| eNOS | endothelial nitric oxide synthase |
| FBS | fetal bovine serum |
| FDA | Food and Drug Administration |
| FSH | follicle stimulating hormone |
| FSR | fractional synthesis rate |
| Gadd45 | Growth Arrest and DNA Damage 45 |
| GAML | GC-associated muscle loss |
| GC | glucocorticoid |
| GCN2 | General Control Nonderepressible 2 |
| GFP | green fluorescent protein |
| GH | growth hormone |
| GLUT | glucose transporter |

|  |  |
| --- | --- |
| GnRH | gonadotropin-releasing hormone |
| GR | glucocorticoid receptor |
| Grb2 | Growth factor receptor-bound protein 2 |
| GRE | glucocorticoid responsive-elements |
| GSK3 | glycogen synthase kinase 3 |
| HAT | histone acetyltransferase |
| HDAC | histone deacetylase |
| HEPES | 4-(2-hydroxyethyl)-1-piperazineethanesulfonate |
| HIF 1a | hypoxia-induced factor 1a |
| HOMA-IR | homeostatic model assessment - insulin resistance |
| HPA | hypothalamic - pituitary - adrenal |
| HS | horse serum |
| HSD | honest significant difference |
| IBMX | 3-isobutyl-1-methylxanthine |
| IGF-1R | IGF-I receptor |
| IGF-I | insulin-like growth factor I |
| IGF2 | insulin-like growth factor 2 |
| IGFBP | IGF-I binding protein |
| IL-1 | interleukin 1 |

|  |  |
| --- | --- |
| IL-1RII | interleukin 1 receptor type II |
| IL-2 | interleukin 2 |
| IRS | insulin receptor substrate |
| IκBα | inhibitory κBα |
| KLF-15 | Krüppel-like factor-15 |
| LC3 | microtubule-associated protein 1 light chain 3 |
| LH | luteinizing hormone |
| LKB1 | Liver Kinase B1 |
| MAFbx | muscle atrophy F-box |
| MAPK | mitogen-activated protein kinases |
| MEF2A | myocyte enhancer factor 2A |
| MNK2 | mitogen-activated protein kinase–interacting kinase  2 |
| MR | mineralocorticoid receptor |
| MRF | muscle regulatory factor |
| Mrf4 | myogenic regulatory factor 4 |
| mTOR | Mechanistic Target of Rapamycin |
| mTORC1 | mTOR complex 1 |
| MuRF1 | muscle RING finger 1 |

|  |  |
| --- | --- |
| Myf | myogenic factor |
| MyHC | myosin heavy chain |
| NEFA | non-esterified fatty acids |
| NFAT | nuclear factor of activated T-cells |
| NMJ | neuro-muscular junction |
| NMR | nuclear magnetic resonance |
| OMIM | Online Mendelian Inheritance in Man |
| ORF | open reading frame |
| p70-S6K | ribosomal protein S6 kinase, 70 kDa |
| p90-RSK | ribosomal protein S6 kinase, 90 kDa |
| PAGE | polyacrylamide gel electrophoresis |
| PBS | phosphate buffered saline |
| PDK1 | 3-phosphoinositide-dependent protein kinase 1 |
| PGC-1 | PPARγ coactivator 1 |
| PI3K | phosphatidylinositol 3-kinase |
| Pitx | paired-like homeodomain |
| PLC | phospholipase C |
| POMC | pro-opiomelanocortin |
| PPAR | peroxisome proliferator-activated receptor |

|  |  |
| --- | --- |
| PPP | picropodophyllin |
| PRAS40 | proline-rich Akt substrate of 40-kDa |
| qRT-PCR | quantitative real-time polymerase chain reaction |
| REDD1 | regulated in development and DNA damage responses-1 |
| RIPA | radioimmunoprecipitation assay |
| SDS | sodium dodecylsulfate |
| SGKL | serum and glucocorticoid-inducible kinase-like kinase |
| Shc | Src homology 2 domain containing |
| Sos | Son of sevenless homolog |
| Sp1 | SV40 promoter-specific 1 |
| STAT | Signal Transducer and Activator of Transcription |
| T | testosterone-only group |
| Tbx | T-box |
| TCA | trichloroacetic acid |
| TCF4 | transcription factor 4 |
| Testo | testosterone |
| TGF-β | transforming growth factor-beta |
| Tp | testosterone propionate |

|  |  |
| --- | --- |
| TSC1 | Tuberous Sclerosis Complex 1 |
| TSC2 | Tuberous Sclerosis Complex 2 |
| ULK1 | UNC-51-like kinase 1 |
| V | Vehicle-only |
| VEGFR | VEGF receptor |

**1. CLINICAL QUESTIONS AND EVIDENCE**

## Cushing’s syndrome

Through the detailed case series written by Harvey Cushing[4], the scientific and medical community became aware of an otherwise rare disease, which bears his name. Unlike the earlier and better-studied deficiencies of the thyroid and pancreas, pituitary defects were more variable in manifestation and therefore harder to unify in a single clinical entity. Even when macroscopic hypertrophy of pituitary was localized to a gland subdomain, it was unclear whether observed pathology could be attributed to a hypersecretion from the hypertrophied sector, or to a deficiency in the neighboring compressed structures. Similarly, pituitary extracts caused multiple, and even opposite effects, in animal models[5], suggestive of a mixture of hormones.

Among 50 cases described by Cushing, about five stood out due to the involvement of other glands. In each of them, and, to a lesser extent, in a few more cases, “hyperadrenalism” was blamed for asthenia, hyperpigmentation of skin, low blood pressure, and hypoglycemia. Histopathology tests localized the adrenal abnormalities to the zona fasciculata of the cortex. Cushing wrote that some of these abnormalities reflect current adrenal hypoactivity, caused by exhaustion after preceding intense stimulation and hyperactivity.

Twenty years later, Cushing narrowed the focus in an updated case series of combined pituitary-adrenal pathology[6]. Cushing noted that basophile adenomata of the pituitary and hypertrophy of the adrenal glands often coexisted. Based on the curative effect of pituitary surgery, he hypothesized that the adrenal defect is secondary to the pituitary abnormality. In turn, he inferred that the adrenal changes mediate the disease phenotype, which includes obesity with

1

ectopic adipose deposits, kyphosis, amenorrhea / impotence, hypertrichosis, lineae atrophicae, fatigability and weakness. Among these disease manifestations, muscle impairment was a serious, if variable, component. Cushing considered intense muscle loss the cause of death for one of these cases.

Cushing’s work did little to elucidate mechanisms leading to the phenotype. The variability in pituitary changes between the cases he described meant that many scientists rejected his hypothesis of pituitary primacy. A group at the Mayo Clinic was actively pursuing the opposite hypothesis, with the adrenal as the primary site of impairment in adrenal-pituitary combined afflictions[7]. On the clinical side, it was noted that some of Cushing’s patients lacked observable pituitary changes. Moreover, some of the Mayo patients were cured by adrenal surgery. From a theoretical perspective, the adrenal hypothesis was more tempting because the adrenal deficiency (termed Addison’s disease) and its reversal by administration of adrenal cortex extracts were better known than pituitary pathology[8].

Today, we know that the truth was more nuanced. Hypersecretion of the adrenal cortex hormones cortisol and / or corticosterone is termed hypercortisolism. One or more clinical signs listed by Cushing (see above) suggest to the practitioner the activation of the hypothalamic - pituitary - adrenal (HPA) axis. If concomitant hypercortisolism is confirmed by an increase of urine free cortisol measurements, or by the effacement of the evening trough in circulating cortisol, there is suspicion for Cushing’s syndrome (CS)[9]. Some hypercortisolism cases, termed pseudo-Cushing’s syndrome, are ascribed to causes outside the HPA axis, such as in depression, morbid obesity, uncontrolled diabetes mellitus, and sleep apnea (reviewed in [10]).

True CS cases are further classified based on the role of the

adrenal-stimulating pituitary hormone corticotropin (adrenocorticotropic hormone; ACTH). In some CS patients, hypercortisolism is paralleled by an increase in ACTH. Their adrenals are usually responsive to further ACTH stimulation tests, indicating that previously intact adrenals underwent hyperplasia in response to a pathological overstimulation with ACTH. When attributed to the pituitary, such ACTH hypersecretion, followed by secondary hypercortisolism, is termed Cushing’s disease (reviewed in [11]). Cushing’s disease remains a staple of physiology textbooks, because it provides an excellent didactic example of a hormone hierarchy.

The remainder of CS cases consists of hypercortisolism in spite of low ACTH.

In primary hypercortisolism, ACTH is typically suppressed by negative feedback.

Adrenal neoplasms are the most frequent cause of primary hypercortisolism. Ectopic or diffuse unregulated sources of ACTH or cortisol may cause hypercortisolism. In recent decades, overdose with synthetic derivatives of cortisol became the most important cause of low-intensity CS (discussed in the next section).

Despite being caused by a diverse set of HPA pathologies, CS is mostly invariable in its ability to cause muscle impairment.

## Glucocorticoid therapy

A series of serendipitous decisions brought impressive knowledge about CS of non-pituitary etiology (reviewed in [12]). First, during World War II, US intelligence learned that Germans were importing large quantities of adrenal glands from neutral Argentina. This reignited US government interest in corticoadrenal research, despite the lackluster results with earlier adrenal extracts. At the end of the war, only a few grams of pure adrenal steroids were manufactured, from endogenous sources and at a high cost. The second opportunity was in the allocation of those scarce steroids. One of them, cortisone, made by Merck, was shared by a few clinical researchers, including Phillip Hench. Hench’s request was based on his previous work on rheumatoid arthritis. He observed that rheumatoid arthritis was alleviated in jaundice, and hypothesized the existence of a steroidal “anti-rheumatoid factor.” Third, Hench’s choice of dose and route elicited an extraordinary reversal in arthritic pain and dysfunction. In 1949, after treating only five patients[13], impressive improvements in those cases triggered redirected corticosteroid research.

Previous work did describe multiple effects for adrenal extracts, but little was known about purer preparations, such as cortisone. In fact, adrenal research was discouraged prior to cortisone purification, because less pure extracts combined antagonistic hormones in variable doses, leading to the impression that they lack a defined pharmacological effect. However, even with purified cortisone, Hench saw a very diverse set of consequences for cortisone administration[14].

First, cortisone’s action on metabolism was accessible even to the less sophisticated clinical measurements used 60 years ago. Patients receiving cortisone gained weight. Chronic cortisone therapy led to accumulation of adipose tissue, often in ectopic locations, such as the interscapular “buffalo hump”. Cortisone also induced hyperglycemia, and subsequent glycosuria. For this reason, cortisone, and its endogenous and synthetic analogs, are grouped in the glucocorticoid (GC) family.

Hench and collaborators hypothesized that cortisone’s protective action is not limited to rheumatoid arthritis. In his 1950 Nobel lecture, Hench envisaged a role for alleviation of most inflammatory diseases. GCs share the ability to reduce inflammation (reviewed in [15, 16]). Some of these anti-inflammatory effects, such as reduction in the number of circulating white blood cells, are ample and robust. The cellular and molecular anti-inflammatory mechanisms are still subject of active research.

The knowledge gap around the anti-inflammatory actions of GCs is in part caused by the immunology progress. Still, some questions remain open, and illustrate the convoluted ways in which GC signals are relayed in the cell. For example, GCs are often acting in a manner shared with all steroids, by binding and activating the glucocorticoid receptor (GR). Activated GR translocates from cytosol to the nucleus, where it dimerizes on specific DNA sequences, termed glucocorticoid responsive-elements (GRE) [17]. The classical effect of the

GRE-GR interaction is increased transcription for neighboring target genes

(transactivation), as it is the case in polymorphonucleate cells with interleukin 1

(IL-1) receptor type II (IL-1RII) [18], a decoy inhibitor of the pro-inflammatory

IL-1. In other circumstances, the activated receptor inhibits transcription directly (transrepression), or by interfering with transcription factors. For example, in human T lymphocytes, GCs inhibit the transcription factor activator protein 1 (AP-1), thus causing a reduction in their ability to synthesize pro-inflammatory interleukin 2 (IL-2)[19]. GCs employ nongenomic mechanisms, such as mRNA stability and enzymatic activity modulations. In airway epithelia, GCs reduce the half-life of the mRNA for interleukin 8 (IL-8), the major chemoattractant for neutrophils[20]. Within minutes, GC administration induces vasodilation, through direct, nongenomic activation of phosphatidylinositol 3-kinase (PI3K) leading to activation of endothelial nitric oxide synthase (eNOS)[21].

Some GC effects may be limited to a range of doses, durations, and frequencies of administration. Moreover, the example of adrenalectomized rats re-supplemented with corticosterone, their most abundant endogenous GC, illustrates how, at times, the same GC can induce or repress the same cellular response, depending on the dose. A 5 mg / kg physiological dose enhances the immune skin delayed-type hypersensitivity, while a pharmacological 40 mg / kg dose yields the more typical GC immunosuppressive behavior [22]. This behavior, suggestive of a U (or inverted U) shaped response curve, is called hormesis or biphasic response, and poses great challenges, both to the investigative scientist and to the clinician attempting to establish a therapeutic regimen.

In 1950, endogenous GCs corticosterone and cortisol, were synthesized at Merck[23], thus lowering the price and creating the opportunity for large-scale trials. The Empire Rheumatism Council organized a randomized trial comparing cortisone with acetylsalicylate, and concluded that there is no benefit in cortisone[24]. While participants receiving cortisone claimed an improvement in subjective well-being, they were afflicted more often with deleterious side effects, including edema and hypertension. In retrospect, a comparison between two palliative symptomatic therapies using cure-indicating outcomes was likely misleading.

By the end of this trial, the chemists were providing more effective and safe synthetic GCs (reviewed in [25]). While synthesizing esters with a longer half-life, Schering chemists introduced a double bond in the A ring of cortisone, thus discovering prednisone, the first widely used oral GC[26]. In addition to an improved ability to induce hyperglycemia, prednisone lost some of the cortisone’s ability to cause edema. NIH researchers synthesized and characterized prednisone’s active metabolite, prednisolone[27]. In a trial of prednisolone versus acetylsalicylate in rheumatoid arthritis, the GC provided better functional protection to the articulations[28].

At Squibb, insertion of a halogen atom was found to abolish GCs ability to

induce hyperglycemia[29, 30]. In 1958, Merck chemists led by Arth modified cortisol with the unsaturated A ring (Δ1), the fluoride addition at position 9α, and with a methyl group on the 16α position to obtain dexamethasone (Dexa) [31, 32]. Dexa is the most effective and specific therapeutic synthetic GC to date, with 170 times higher ability to inhibit the immune reaction to subcutaneous foreign bodies (granuloma) compared to cortisol.

Dexa is unable to cause edema and electrolyte imbalance. Nevertheless, Dexa is 52 times more potent in suppressing endogenous GC secretion, and 35 times more potent in causing hyperglycemia[33, 34]. In animals, Dexa is 160 times more potent than cortisol in the inhibition of granuloma test of immune function[35]. These findings suggest that anti-inflammatory and hyperglycemic actions are intermediated by the same specific, Dexa-sensitive receptor, whereas electrolyte changes are caused by cortisol through a different pathway. Further supporting the idea of a unique Dexa receptor, efforts to synthesize steroids with anti-inflammatory action that do not interfere with metabolism have failed. Compounds such as A276575[36] and RU 24858[37] did not reach the human studies stage. Mapracorat[38] did nor progress beyond phase II clinical trials. Therefore, clinicians prescribing GCs in these five decades had to balance therapeutic benefit with metabolic side-effects.

Edema is an example of non-specific GC effect, caused by a less typical interaction of the hormone with the mineralocorticoid receptor (MR). The GC family spans tens of active principles and thousands of formulations, from weak

GCs with lower specificity, such as cortisone, to strong, specific GCs, such as Dexa. The list of Food and Drug Administration (FDA)-approved indications for cortisone, Dexa, and prednisone is often changed, as better, more specific drugs are developed, and more precautions are added[39, 40, 41].

The trivial case for using GC therapy is in hormone replacement, such as in adrenocortical insufficiency (reviewed in [42]). GC therapy is suitable for acute immune or allergic conditions, such as seasonal rhinitis (reviewed in [42]). GCs are relatively safe in topical applications in dermatological conditions

(pemphigus, psoriasis, most types of dermatitis; reviewed in [43]). Similarly, GCs are commonly used in eye inflammatory conditions[44, 45], such as diffuse posterior uveitis and optical neuritis.

As envisaged by Hench in his Nobel Lecture, GCs do not address disease causes and are recommended for temporary respite. For many autoimmune diseases, there are more specific therapeutic alternatives, often addressing the cause of disease. In addition to the glucose metabolism disturbance, long term GC administration causes osteoporosis and muscle loss. On the balance of benefits and drawbacks, GCs are recommended for life-threatening or impairing immune reactions, such as in polymyositis (reviewed in [46]), severe sarcoidosis (reviewed in [47]), and disseminated pulmonary tuberculosis. Based on their ability to lower white blood cell count, GC are an important adjuvant in the palliative and even etiologic treatment of leukemias and lymphomas[48, 49, 50].

Even in chronic diseases, GCs are recommended for short-term alleviation of exacerbations. Short-term GC therapy is recommended for rheumatoid arthritis, gouty arthritis, psoriatic arthritis, ankylosing spondylitis, asthma[51, 52], ulcerative colitis[53, 54], and idiopathic nephrotic syndrome[55].

GCs have been used off-label in many other diseases. Most commonly, GCs are perceived by physicians as a fallback therapeutic alternative for cerebral hypertensive conditions, despite sparse evidence for efficacy in specific conditions. Small trials suggest GCs reduce vasogenic cerebral edema[56] and prevent acute mountain sickness[57], while systematic reviews suggest they might in fact worsen outcomes for acute brain trauma victims[58]. A similar, paradoxical situation is seen in ongoing clinical research. As of 2015, the patent-free status of the GCs discourages trials for new indications, while their de facto standard-of-care status makes them a common comparator in clinical trials. The National Cancer Institute sponsors 311 ongoing clinical studies employing Dexa, mainly in the standard-of-care arm, thus providing a plethora of data that have been, and may still be, misconstrued as support for the use of GCs. Every day practice may drift further apart for the officially sanctioned label, thus providing new opportunities for unjustifiable overdose.

Due their widespread use, GCs are likely to cause covert iatrogenic CS in a large population, impairing muscle and quality of life to a certain and understudied degree.

## Hypercortisolism-induced muscle loss

Primary and secondary endogenous hypercortisolism are rare diseases (1-2 cases per million and year each[59]), despite a recent boost from incidental findings during imaging tests for other needs. The symptomatology is non-specific, meaning that, even today in the developed world, an average of 6 years pass from signs onset until diagnosis is made and treatment is initiated[60]. It is a life-threatening disease, with untreated patients having a median survival rate of 5 years after diagnosis[61]. Some of the changes occurring in Cushing’s disease are irreversible, especially at the level of brain, bone, adipose tissue, and liver levels (reviewed in [62]). Even after surgical adjustments of the hyperactive pituitary, the quality of life for CS patients lags behind that of the unaffected population.

At presentation, about two thirds of Cushing’s syndrome cases present with muscular complaints, with similar incidence among pituitary and adrenal conditions[63]. Among patients suspected of endogenous CS, one fifth are referred to the endocrinologist due to muscle weakness[64]. Two-fifths of those whose endogenous hypercortisolism is successfully corrected by surgery still complain of fatigue[65].

On the other hand, therapy-induced (iatrogenic) CS is common. The glut of GC indications and off-label uses makes them some of the most used drugs in the developed countries. Every year, about 1% of the Americans and British receive some form of GC[66, 67]. GCs are likely even more often prescribed in the developing world, due to affordability and lack of alternatives, poor access to health care notwithstanding. Dexa and cortisol are the only drugs listed five times in the World Health Organization’s List of Essential Medicines[68].

In most cases, the cause of iatrogenic CS can be identified by careful history taking and medication reviews. However, an increasing number of cases are not as easily diagnosed, because the excess GC is not from prescribed medicine. In United States, FDA approved in 1979 over-the-counter sale of 0.5% hydrocortisone cream for itching and minor skin inflammation. In 1990, 1% hydrocortisone creams were also permitted[69]. Where regulated, over-the-counter GC creams rarely cause CS on their own, but may lower the threshold for CS in patients who are also prescribed oral GC. Unregulated, mislabeled, overdosing GC creams sold as skin-bleaching products pose a great CS risk to patients from ethnic groups with darker skin. Half of the respondents in a Nigeria poll admitted using GC-based skin bleaching products[70]. In 2015, the Ivory Coast government made illegal skin bleaching products, due to worries about GC overdose side effects[71]. The side effects of skin bleaching are well recognized by the sub-Saharan medical community. Paradoxically, CS caused by bleaching products may be less identifiable to practitioners who care for the African diaspora in the developed world, where bleaching is more frequent, due to improved financial access and social pressures[72, 73].

Other, less frequent causes of iatrogenic CS include the interaction between low dose GC therapy and cytochrome P450 3A4 inhibitors, such as the antiretroviral ritonavir[74]. Other steroid drugs may interact with GR and cause CS when overdosed, as it is the case with the synthetic progestin megestrol

acetate[75].

Due to its insidious and erratic symptomatology, iatrogenic CS is often diagnosed years after onset or completely unrecognized[60]. The incidence of iatrogenic CS is difficult to estimate, because there is no reporting requirement. In the developed world, iatrogenic CS could be as frequent as one case per thousand and year[76].

Signs of iatrogenic CS are as varied as those of Cushing’s disease. In a cohort of patients receiving for three months more than 0*.*4 mg/(kg d) prednisone, the most common signs were development of ectopic adipose deposits (50%), hyperphagia (47%), and muscle cramps (32%) [77]. In the same cohort, 15% complained of muscle weakness. Patients stated that the most distressing signs of hypercortisolism were, in order, body shape changes, neuropsychiatric disorders, muscle cramps, and hand tremor. Mastaglia estimated that, in 1982, the most common cause of iatrogenic muscle weakness was caused by GC[78].

There are differences between GC-induced cardiovascular changes, depending on the nature of the GC. Endogenous GCs, such as cortisol, have hypertensive effects, while some synthetic GCs, Dexa included, lack such non-specific MR-dependent action. Nevertheless, excess exogenous and endogenous GC cause the same disabling effects on muscle[79], indicating that muscle damage is mediated by GR. GCs do differ quantitatively in their ability to cause myopathy. Myopathy is invariably induced in two weeks by either 0*.*2 mg/(kg d) Dexa[80], or by 0*.*5 mg/(kg d) prednisone[81]. Based on other metabolic effects, it is likely that the catabolic potency ratio is even more tilted towards Dexa than the referenced studies indicate. More detailed human pharmacodynamics studies are needed.

In their 1958 case series, Muller and Kugelberg were the first to describe muscle changes associated with long-term Cushing’s disease[82]. In their mixed, primary and secondary, endogenous hypercortisolic cohort, they found that complaints of muscle weakness were primarily focused on the thigh. Objective loss of muscle force was correlated with histopathological changes indicative of a muscle fiber defect, such as degenerated fibers, at times hyalinized or with loss of striation, muscle replacement with fat and connective tissue, and rare hypertrophic fibers. Through electromyography, they established that the number of motor units is unaffected. Together with lack of changes in reflexes, their work negated a neurological component of CS. Muller and Kugelberg noted faster extinction of the action potential, which may be caused by a reduction in the number of fibers, or by fiber atrophy[83]. Based on the evidence that GC is a muscle fiber disease, they coined the phrase “steroid myopathy.” Similar electromyography changes are induced by long-term GC therapy[84], making some authors reserve the term “steroid myopathy” to muscle complaints of iatrogenic etiology. In 1966, D’Agostino and Chiga, confirming histological fiber changes in a rabbit model of iatrogenic CS, formulated the more precise, yet less commonly used “glucocorticoid myopathy”[85]. Owing to the fact that steroid myopathy is not a standalone disease or syndrome, terminology has never been standardized. In the present work, where animal models are examined, the condition of interest will be termed GC-associated muscle loss (GAML).

In exogenous CS, GC excess can be better quantified. In a population with neurological maladies receiving long-term Dexa, the threshold for manifest steroid myopathy appears to be 50 µg/(kg d)[86]. However, the most significant predictor of clinical GAML is total dose[80, 87]. When GAML develops, the amplitude of electromyography changes (that is, the reduction in action potential duration) is proportional with the total GC dose[88]. These findings imply that steroid myopathy can be induced in shorter periods, if the GC dose is extremely high. Foye and colleagues drew a distinction between “classical” chronic steroid myopathy, induced “within weeks to years,” and acute steroid myopathy, induced in 5-7 days of high-dose GC[89]. However, their description of the two forms of GAML is almost identical, suggesting that the two clinical entities are largely overlapping.

In a comparative study of patients receiving GC therapy for asthma, half of the patients receiving more than 0*.*2 mg/(kg d) prednisone exhibited a reduction in hip flexor strength of 2 SD or more, compared with health age- and sex-matched controls[81]. In a study of adults with brain or spine cancer, 60% of the participants experienced loss of iliopsoas muscle force in response to GC therapy for cerebral edema[80]. In a small cohort, 6 months of 0*.*16 mg/(kg d) prednisone treatment was associated with a 20% reduction in thigh muscle force, compared to healthy controls[90]. In a post-hoc analysis of a chronic obstructive pulmonary disease (COPD) trial, the placebo arm was stratified in GC-treated and GC-naive groups[91]. The maximal inspiratory mouth pressure, a proxy measurement for respiratory muscle strength, was significantly better maintained over the 8 weeks of the trial in the GC-naive, compared to GC-treated participants. Such findings suggest that GC-induced weakness is caused by an objective muscle disorder, and negate the alternative, neuropsychiatric etiology.

Another investigative direction in the study of GC-induced muscle weakness focused on muscle mass and volume. Although correlated, muscle force and muscle mass are not completely reflecting each other. The most accessible proxy measurements of muscle mass, such as mid upper-arm or thigh circumference, are not sensitive enough in monitoring GC-induced muscle loss, even after subtracting skin fold, because GC stimulate intramuscular adipose deposits[92]. The advent of modern imaging allowed non-invasive muscle measurements. Chronic prednisone administration causes a 20% reduction in mid-thigh muscle area measured by computed tomography, and a 36% increase in the ratio of fat-to-muscle areas (CT)[93]. Psoas muscle area and density, measured by computed tomography, are inversely correlated with GC levels indicated by 24-hour urine cortisol (24HUC)[94].

In an early study of chronic hypercortisolism, it was found that all types of fibers are affected by GC[95]. In more recent investigations, a type-specific effect was described. Women with CS have an increased proportion of type IIx (fast twitch, glycolytic) and a lower proportion of type I (slow twitch, oxidative) fibers in their vastus lateralis muscles[96]. Similar histological findings were made in renal transplant patients receiving 25 mg/(kg d) prednisone over three months[97]. In the latter, GC caused an increase in the cross-sectional area (CSA) of the type I and IIa (slow twitch, oxidative / glycolytic) fibers. Gains in the ratio fast-to-slow twitch fibers are associated with insulin resistance[98]. Diameter increases in spite of loss of function and protein content have been explained by a disorganized intracellular structure. A more practical consequence of these findings is the dereliction of CSA measurements in GAML.

A set of muscle mononucleate cells, expressing the paired-box transcription factor Pax7, are presumed to support muscle development and regeneration, and are termed satellite cells (reviewed in [99]). There are no definitive studies describing the effect of GC in human satellite cells. Some or all satellite cells may be activated to proliferate, thus becoming myoblasts. Many in vitro assays use proliferating cells from human muscle, at times assumed to be myoblasts. These human “myoblasts” do not proliferate in the absence of at least 1 µM ([100], and personal observation; data not shown). For comparison, maximum normal concentration of endogenous cortisol in humans is 0*.*78 µM[101], that is, tens of times less potent. Therefore, it is impossible to conceive an experiment where these human myoblasts are subjected to meaningful manipulations of GC concentration. Moreover, it appears that GCs are vital for human muscle development and maintenance. This implies that cell lines that do not require GC, such as those surviving in serum-free media, may be less accurate models of muscle.

There are no published cases of increase in circulating myoglobin or creatine kinase in response to GC monotherapy, or as a consequence of Cushing’s disease. The absence of such intramuscular protein from the blood flow suggests GC do not cause rhabdomyolysis, that is, loss of muscle through uncontrolled rapid membrane leakage.

From its first trial, GC therapy ability to induce a negative nitrogen balance, through an ample increase in urinary creatine and creatinine, was interpreted as evidence for stimulation of tissue protein breakdown[102]. As little as 20 µg/kg cortisol infused over 8 hours increases by a quarter the rate of appearance of leucine into the bloodstream, suggestive of proteolysis upregulation[103]. Leucine’s rate of appearance is even higher when the GC-induced hyperinsulinemia is prevented, indicating that whole-body experiments do not capture the amplitude of the GC-induced proteolysis[104]. More modern mass spectrometric methods revealed that a single dose of 1 mg/kg prednisolone cause increases in all blood amino acids, presumably due to mobilization from muscle sources[105]. The same acute treatment causes an increase in 3-methylhistidine (3MH), a degradation product specific to muscle actin and myosin[106]. Similar increases in 3MH are seen with control diet in chronic GC excess of endogenous or exogenous nature[95]. These findings demonstrate that GC-induced loss of muscle mass is mediated by stimulation of protein degradation.

The last three decades brought a better understanding of protein degradative pathways and of muscle atrophy. Two major proteolytic systems, the proteasome - ubiquitin system and the autophagosome (discussed in later sections), have been discovered and dissected. Unfortunately, only one published trial investigated the action of GC in human muscle biopsies, at a molecular level. It failed to find a significant change in mRNA of ubiquitin and the C3 subunit of the proteasome[107]. The result is unsurprising, given that the control of the proteasome system may be exercised in other, unexplored ways. In animal models, the genes most correlated with muscle loss, including GAML, are two E3 ubiquitin ligases, muscle atrophy F-box (MAFbx; gene known as Fbxo32) and muscle RING finger 1 (MuRF1; gene known as Trim63), but no published studies confirm or refute their modulation in humans (reviewed in [108]).

Recently, pharmacological inhibitors of the proteasome became widely available. The first proteasome inhibitor, bortezomib, is recommended by the FDA for multiple myeloma and mantle cell lymphoma[109]. The second generation, irreversible proteasome blocker carfilzomib is also approved for advanced myeloma therapy[110]. In the light of data from the animal models of muscle loss, these drugs should have been useful in cachexia, but, to date, no human trials investigated their ability to prevent muscle atrophy.

There are no trials comparing GC with the combination (GC + bortezomib). However, an indirect comparison can be made. In a trial for multiple myeloma, fatigue was a complaint of 32% of the participants receiving 40 mg Dexa, compared to 42% for bortezomib[111]. In another trial, addition of 20 mg Dexa to bortezomib lowered the rate of fatigue from 57% to 25%[112]. Taking into account the large variability between trials and the use of an atypical population, the comparison is of marginal use, but it does not appear that the combination (Dexa + bortezomib) is more muscle protective than Dexa alone. Clinical studies directly addressing this comparison in patients prescribed high-dose GC are recommended, given that the most commonly accepted hypothesis centers on the proteasome as main effector of GC-induced muscle loss. It is still more likely to find that bortezomib provides muscle protection. Proving a beneficial action of bortezomib in co-administration with GC will have major practice implications. Even proving the opposite, that bortezomib has no protective action, will be very valuable in better understanding and eventually preventing GC-induced muscle

loss.

The inhibition of the other proteolytic system, the autophagosome, is also the focus of clinical studies. Starting with the inexpensive antimalarials chloroquine and hydroxychloroquine, autophagosome inhibitors are now the focus of phase II clinical studies in many cancers[113]. Interestingly, hydroxychloroquine is also recommended for rheumatoid arthritis, where it may be prescribed for up to six months[114]. Chronic hydroxychloroquine therapy is known to induce muscle weakness and sporadic myopathy, through a distinct, vacuolar mechanism. The hydroxychloroquine-induced myopathy is associated with an increase in autophagosome markers in muscle, demonstrating the importance of autophagosome in muscle regulation[115]. In two separate case reports, co-administration of prednisone and hydroxychloroquine led to vacuolar myopathy, which could be caused by the choice of doses, or could be indicative of true epistasis[116, 117]. Potential benefits of anti-lysosome co-therapy for the atrophying muscle remain the subject of speculation.

Another putative parallel mechanism for GC-induced loss of muscle is downregulation of protein synthesis. Few human trials measured directly the effect of GC on protein synthesis in healthy volunteers. Brillion and colleagues[104] found that an 80 mg cortisol infusion over 13 hours led to 8% increase in non-oxidative leucine uptake, indicating an upregulation of protein synthesis. However, using a 200 mg cortisol infusion in the same protocol failed to cause a detectable change in protein synthesis compared to placebo, suggesting hormesis may confound experiments. Moreover, this finding was never replicated. Löfberg and colleagues[107] found that three days of 65 mg / day prednisolone caused a non-significant 21% increase in protein synthesis rate and a statistically significant 52% increase in the rate of protein degradation, based on the difference between arterial and venous levels of tritiated phenylalanine at leg level. Short and colleagues employed fractional synthesis rate (FSR), which describes the time rate of enrichment in muscle tracer, normalized to the circulating tracer concentration. They concluded repeatedly that, in leg muscles, 35 mg / day prednisone for 6 days “has no effect on [...] muscle protein metabolism or muscle function”[118, 119]. Some of these studies may have been underpowered (sample size n = 6-7) or may be troubled by the use of a small dose, but their validity is confirmed by the fact that, in each case, the expected hyperglycemic response to GC was observed.

The hypothesis that GC cause muscle loss by inhibition of protein synthesis is still debated, due to a plethora of indirect evidence. In Löfberg’s study, biopsies revealed a prednisolone-induced loss of muscle polyribosomes, interpreted as evidence for decrease in protein synthesis rate. Even in studies where GC failed to elicit reductions in protein synthesis, they inhibited translation-stimulating signals in muscle from anabolic factors such as insulin[120], branched chain amino acids[121], and exercise[122].

At a molecular level, it appears that Dexa inhibits anabolic signals centered on the Akt / mechanistic target of rapamycin (mTOR) axis. More evidence has been obtained from animal models (discussed in a dedicated section). One study on humans described how Dexa inhibits branched chain amino acids’ ability to induce phosphorylation of eukaryotic translation initiation factor 4E (eIF4E) binding protein (4E-BP), isoform 1, and p70-S6K[123]. Dexa lacked similar action on another translation regulator, eIF2α.

In addition to GC excess, muscle weakness is caused by GC withdrawal[124], and by GC deficiency, illustrated by the Addisonian crisis[125]. In both hypercortisolism and hypocortisolism, effects on human muscle remain understudied. Human studies concord that GC-induced loss of muscle force is an objective finding caused by an increased proteolytic activity. Indirect evidence indicates that human GAML is associated with changes in protein synthesis. In the absence of other proven mitigating interventions, current guidelines suggest GC discontinuation if myopathy develops. Animal models have been essential for the study of GC-induced muscle loss, although they have been confounded by hormesis (discussed in future sections).

In conclusion, CS of various etiologies leads to an increase in muscle protein catabolism, and a reduction in muscle protein synthesis.

## Muscle protection with androgen therapy

A series of historical circumstances brought anabolic androgenic steroids (AAS) in the attention of clinicians treating hypercortisolism in muscle. The same circumstances meant that utility of AAS therapy in steroid myopathy has never been fully explored.

Male hormones have been considered an efficacious anabolic therapy long before they were purified and tested. The effects of male castration, such as reductions in aggressiveness and muscle force, were discovered independently by many human civilizations, starting more than three thousand years ago. Castration is omnipresent in ancient mythology, and, more mundanely, in primitive farming. For almost as long, people perceived testis ingestion as a reversal of castration, thought to improve muscle force. Such perceptions were caused by the placebo effect alone, given that this testis active principle is almost completely degraded by liver.

Testis extract benefits received more attention starting around 1889, when Brown-Séquard published his theory about rejuvenating abilities of sperm. He thought that loss of sperm during aging or masturbation causes degradation in muscle and brain performance, and hypothesized that chemicals from sperm may pass into blood where they have “a most-essential use in giving strength to the nervous system and to other parts.” Consequently, he injected himself with a combination of sperm and testis extracts, which led to self-reported improvements in physical and intellectual abilities[126]. He describes how, at the age of 72, a single injection enables him stand for hours, or write longer scientific papers. Later on, he describes how testis extracts appeared to alleviate “serious affections of any kind,” including cachexia, pulmonary tuberculosis, cancer and leprosy ulcers[127]. Because the active principle in testis is made as needed, rather than stored in high-concentration depots, it is now obvious that these observations were the product of preconception.

The cultural context in which Brown-Séquard worked introduced multiple biases in his experiments and conclusions. His mistaken theses were constrained into rather low-quality experiments, which luckily provided useful, testable, and eventually proven scientific hypotheses. First, the logical conclusion for

Brown-Séquard’s theory would have been endorsement for semen therapy. Instead, due to the semen taboo, Brown-Séquard and his disciples resorted to surrogate interventions, such as vasectomy, believed to preserve sperm in the body, and injections with testis extracts. The introduction of injections gave a new lease of life to the therapeutic use of organ extracts, called “organotherapy,” which had been banished from the British Pharmacopoeia in 1788 after failing the test of oral administration. Some organotherapies were shams or even harmful. Yet a few of them provided evidence that specific parts of the body store or release into the blood stream chemicals, which subsequently induce changes in other specific parts of the body. This conjecture led the discovery of endocrine glands and the establishment of endocrinology as a science. In fact, androgen organotherapy provided the pattern to GC discovery.

Second, the Victorian era is an age of body rediscovery. Georgian pastimes, such as cock fighting, horse racing, or cricket, are replaced by more muscular sports, such as football, rugby, gymnastics, and swimming. Bodybuilding becomes fashionable, with the first professional competition selling out Royal Albert Hall in 1901. Brown-Séquard’s promise of muscle without effort makes testis organotherapy a widespread, well-earning business. When Voronoff is barred from practicing in Paris and judged as fraudulent by the Royal Society of

Medicine, he takes his testis transplant business to Algiers, where he receives patients from all over the world (reviewed in [128]). Private sponsorship led to investment in androgen research, but with a focus on commercial rather than

clinical efficacy.

Finally, Brown-Séquard’s era tolerated unscientific theories, which ignored the physical and intellectual ability of women. Brown-Séquard claimed that ovary extracts provide some benefits, but with “less power” than testis extracts[127]. Such conclusions stemmed from cultural biases rather than comparative experiments. In 1849, Berthold showed that, through testis implants, roosters regain male characteristics they lost through castration, such as aggressiveness, libido, and larger combs[129]. With maintenance of secondary sex characteristics as its sole ability, Berthold’s secreted agent was therefore androgenic. In contrast, Brown-Séquard claimed that his extract increases muscle force, without mentioning any virilizing side effects. Moreover, in 1935, Kochakian proved that urine-extracted “male hormone” stimulates muscle accretion in castrated dogs, that is, that it is anabolic[130]. While ultimately proven correct, the idea that “male hormones” were simultaneously androgenic, anabolic, and ergogenic was based on a cultural construct that confounded manliness and physical force, rather than the product of evidence.

The belief in a male-secreted ergogenic substance inspired many commercial enterprises to sponsor research in male endocrinology, through the decades where the evidence was confined to changes in the combs of roosters. These dark ages end in 1927, when McGee and Koch extract a lipophilic virilizing mixture from rooster testis[131, 132]. A pure and even more androgenic chemical is extracted in 1935 from bull testis by Laqueur, working for Organon[133]. Laqueur names his discovery testosterone (Testo). Three months later, Butenandt and Ruzicka, sponsored by Schering and Ciba respectively, announced the development of manufacturing methods for synthetic testosterone, an achievement that brought them the 1939 Nobel Chemistry Prize (reviewed in [134]). At the University of Chicago, Kenyon tests Testo on four eunuchoid patients of testicular and pituitary etiology. Daily injections of 25 mg testosterone propionate (Tp) cause an doubling in prostate and penis size[135] after less than two weeks, thus establishing the efficacy of Testo replacement therapy in men with pathological decreases in circulating Testo. Except for a few, narrow exceptions, this population was and remains the only generally accepted, FDA-approved indication for Testo therapy[136, 137, 138]. Recent Testo preparations are still recommended for some breast cancers, but this indication is limited to a few, unpredictable, cases.

Due to manufacturing costs, limited commercial target, and governments’ lack of interest, Testo therapy traversed a very long experimental stage, which could easily be called “the second dark age of androgens.” Only in 1953, FDA gives its first approval for an androgenic therapy, a Testo enanthate injection. Then, as now, FDA’s approval was based on Testo ability to restore normal levels of androgens, rather than other, more functional or curative, outcome[137]. However, in 18 years of life as experimental drugs, androgenic steroids have been trialed in diverse diseases, including male functional impotence[139], unwanted lactation[140], uterine bleeding and dysmenorrhea[141], or osteoporosis[142]. These early studies share the extremely small sample size, and the scarcity of controls, blinding, and objective outcomes. For example, a study found that 14-35 injections of Tp (cumulative dose 255-455 mg) caused an improvement of acne in half of the male participants[143]. Such findings are at odds with more modern trials, where weekly i.m. androgen injection lead to an increase in absolute risk of acne by 15%, in healthy males[144], and are possibly explained by the variability in the androgen arm, small sample size (n = 12), lack of blinding, and early stopping in the placebo arm. Nevertheless, these trials are, in many cases, the only source of information about the action of Testo in the normogonadal population. For example, early trials of oral methyltestosterone revealed its hepatic toxicity, with the effect that, 50 years later, the development of oral androgenic therapies is still discouraged.

The second dark age of Testo was a time of limited knowledge, and even more limited adherence to the principles of clinical research. Yet in these years, androgenic steroids first gained their reputation as ergogenics. Kenyon noted in his studies on eunuchoid men that Testo injections helped them gain weight through protein accretion, as demonstrated by a reduction in urinary nitrogen. Other trials evidenced benefits from androgenic therapy in muscle-depleting conditions, including thyrotoxic myopathy[145] and muscular dystrophy[146]. By 1940, Kenyon confirmed that Tp caused nitrogen retention, caused by increased protein accretion, even in healthy men and women[147]. Interestingly, in 1942, Samuels and colleagues state that Testo does not change grip strength in healthy males[148]. According to a meta-analysis[149] and my literature search, no other test of androgens’ effect on muscle strength is published until 1968. Despite the lack of evidence, androgens are used as ergogenics in healthy people, starting with Olympic athletes around 1954[150].

As exemplified by the ergogenic hypothesis, benefits of androgen therapy on men with Testo deficiency have been extrapolated by clinicians and theoreticians to other muscle-depleting conditions, and even to healthy humans. Naturally, one of the conditions associated with loss of muscle mass that clinicians hoped to improve was hypercortisolism. In 1941, Albright showed that the newly discovered Tp, in 25 mg daily injections, was better than estradiol benzoate, progesterone, or vitamin D in restoring nitrogen balance in three cases of Cushing’s disease[151].

Similarly, in 1950, the Mayo Clinic team who discovered cortisone remarked that, in one case, 25 mg Tp daily injections reduced urinary nitrogen losses caused by 200 mg cortisone administration[14]. Some of the aforementioned researchers publish similar case reports, sharing the small sample size and the use of surrogate outcomes[152]. These shortcomings do not prevent each investigator from subjective claims of improvements in physical function.

During the 1950’s, AAS became part of the standard of care for endogenous hypercortisolism during the gap between diagnosis and curative surgery. However, this gap narrowed to a few weeks, due to improvements in differential diagnosis. Development of accurate cortisol assays allowed the measurement of its changes in response to Dexa, thus discriminating conditions where feedback mechanisms fail (mainly endocrine neoplasms) from cortisol-stimulating non-endocrine conditions. ACTH assays differentiate ACTH-independent cases (typically adrenal tumors) from the ACTH-dependent ones (usually localized in the pituitary). Modern imaging, including computed tomography of the adrenal and magnetic resonance imaging of the pituitary, identify the target of surgery. AAS therapy is now confined to inoperable cases, including unidentified ectopic sources of ACTH or cortisone. Even these cases benefit from more targeted interventions (see1).

Similarly, the opportunities for AAS as adjuvant to GC therapy are very limited. Many of the diseases previously treated by high-dose GC are now treated with more specific drugs. As practitioners became more accustomed with the risks of GC therapy, doses and durations were reduced. With the exception of life-threatening conditions, typical GC prescriptions switched to lower-potency compounds, such as prednisone or even cortisol. In particular, practitioners became well aware of the issues of GC withdrawal syndrome, where adrenal

|  |  |
| --- | --- |
| Class | Medications |
| ACTH  inhibitors | * Subtype 5 somatostatin receptor agonists: pasireotide (FDA-approved)[153] * Dopamine D2 receptor blockers:   cabergoline |
| 11-β hydroxylase inhibitors | Metyrapone, mitotane, ketoconazole. |
| Inhibitor of 3 βhydroxysteroid dehydrogenase | Trilostane (EMA-approved, FDA-withdrawn)[154] |
| Inhibitor of the cholesterol side-chain cleavage enzyme | Aminoglutethimide |
| GR antagonist | Mifepristone (FDA-approved)[155] |

Table 1: Pharmacological agents used in Cushing’s disease of unidentified ectopic, or diffuse localization (reviewed in [1, 2])

atrophy is aggravated by some other, still undiscovered, component[124]. By mid-1970’s, it became common advice that “prescriptions for [glucocorticoid] steroids should not be refillable”[156]. By the time modern trials with AAS began, the incidence of overt hypercortisolism decreased significantly. Despite a potential epidemic of covert hypercortisolism, with deleterious effects of life quality and expectancy, the interest for studies on hypercortisolism has largely waned. Clinical studies investigating the benefits of AAS in hypercortisolism are scarce and small-scale. For example, there are no significant-size clinical studies analyzing the effect of AAS on the muscle strength of the endogenous CS patient.

An unblinded trial observed AAS-induced increases in lean body mass and appendicular muscle mass, in men already receiving an average of 6 mg prednisone a day over 9 years[157]. Another, randomized, blinded, placebo-controlled trial by Crawford and colleagues tested the benefits of testosterone or nandrolone decanoate as an adjuvant to chronic GC therapy for diverse pathologies[158]. The exposure to GC was an average of 12 mg prednisone a day, over more than 8 years, and was already causing osteopenia, hyperlipidemia, hypercholesterolemia, and a reduction in quality of life compared to historical controls[159]. Such side effects could arguably be considered evidence for mild iatrogenic CS. After six months of 200 mg testosterone injections every other week, the AAS group had higher bone density, muscle mass and strength, and a better quality of life, compared to the placebo group. To date, Crawford’s study is the best evidence for effectiveness of AAS as adjuvant in GC therapy.

In a subset of CS patients, androgen administration improves muscle mass, and, presumably, quality of life.

## Hypercortisolism-induced changes in endogenous androgens levels

Muscle wasting is more common in males than in females with hypercortisolism[160]. The interaction of GC with endogenous AAS distinguishes male from female hypercortisolism, to the point that the two syndromes are

qualitatively different.

The period when AAS were frequently used as therapy of CS or as adjuvant to high-dose GC antedates modern molecular biology and genomics. Therefore, there are no published human trials to describe in molecular terms the interaction of GCs and AAS at muscle level. Most of our knowledge is derived from animal models (discussed later). On the other hand, clinical observational studies of circulating biomarkers remain common, and reveal an interesting interaction between the two classes of steroids. More specifically, in many cases, hypercortisolism suppresses endogenous AAS. Because loss of endogenous AAS, also termed hypoandrogenism, is associated with loss of muscle mass and strength[161, 162], an AAS replacement strategy in hypoandrogenic hypercortisolism may be beneficial for the muscle.

A series of trials observed the effect of short-term (hours or days) hypercortisolism in healthy volunteers. Experimental acute hypercortisolism represses circulating levels of Testo, in a reversible manner, in males and, to a lesser degree, in females [163, 164]. The mechanisms through which hypercortisolism causes hypoandrogenism are still to be elucidated. Some studies suggest that acute hypercortisolism downregulates the pituitary-secreted, T-upregulating, luteinizing hormone (LH)[165, 166, 167]. Others counter that GC induce hypoandrogenism even when LH is unchanged[168]. Another hypothesis is that the negative feedback loop repressing ACTH in hypercortisolism has a side effect of androgen suppression[169]. A few groups have even hypothesized the existence of another, still unknown hormone, synthesized from ACTH precursor, pro-opiomelanocortin (POMC), with the ability to stimulate androstenedione synthesis and secretion[170, 171]. Certainly, GC-induced repression of POMC should also repress this unknown androgen-stimulating hormone, but its existence was never proven.

In males, chronic hypercortisolism is also associated with hypoandrogenism. Long term prednisone therapy reduces circulating Testo levels[172]. Similar observations have been made in endogenous CS, where exposure is longer and, depending on etiology, ACTH is increased or decreased compared to normal. Some studies found that, in CS, LH and another gonad-stimulating pituitary hormone, follicle stimulating hormone (FSH) are lower than normal[173]. This has been explained as a CS-associated pituitary defect, with loss of LH response to stimulation by its hypothalamic regulator, gonadotropin-releasing hormone (GnRH)[173, 174]. Alternatively, others concluded that hypercortisolism impairs hypothalamic GnRH secretion[175]. Finally, a small study found that male asthma patients receiving long-term prednisone have lower circulating Testo levels despite increases in LH and FSH, and concluded that prednisone has an direct inhibitory action on the testes[176]. Despite disagreeing on the mechanism, all these studies agree that chronic hypercortisolism represses testicular androgen secretion.

AAS therapy does not change circulating cortisol levels[177], suggesting that a reverse effect probably does not exist. Similarly, it is possible that the direct effect of GC on AAS is only an artifact caused by pathological and pharmacological doses.

In both sexes, the most concentrated circulating steroids are dehydroepiandrosterone (DHEA) and its ester, DHEA sulfate (DHEAS), which originate from the adrenal and, to a lesser degree, from gonads. Their most important role appears to be that of precursors for synthesis, in glands and peripheral tissue, of androgens and estrogens. DHEA has some affinity for the AR, which suggested it may be an AAS. Recent studies indicate that, in human female tissue, DHEA may in fact be a partial agonist, hindering the action of T[178]. DHEA and DHEAS, now termed adrenal androgen precursors (AAP), are upregulated by ACTH, through increased synthesis of DHEA in the adrenal and rapid bidirectional interconversion[179, 180]. Therefore, Cushing’s disease and other conditions associated with increases in ACTH will present with increases in AAPs, while primary hypercortisolism will be associated with ACTH repression and consequent AAP decrease[181, 182, 170, 183]. Both types of hypercortisolism manifest GAML, despite opposite effects on AAPs, suggesting that AAPs changes are not mediating GAML.

In adult women, the regulation of AAS is more complex. During reproductive age and a few years afterwards, the main source of androgenic stimulation is the ovary[184], where Testo is an intermediate product in the synthesis of estrogens (reviewed in [185]). A feedback loop links LH and estrogens levels, with LH stimulating synthesis and secretion of estrogens from the developing and atretic follicles[186]. The reverse link is more complex, with estrogens inhibiting LH for most of the menstrual cycle[187], with the possible exception of ovulation. In the direct link, LH must stimulate ovarian Testo synthesis, but a reverse link, where Testo directly inhibits LH, is absent in women[188, 189]. Although measurement methods and normal ranges are still to be perfected, it appears that circulating Testo level in women are reflecting the menstruation-related cyclical interplay of estrogen and LH, rather than being independently controlled[190, 191].

This sexual dimorphism differentiates male and female AAS response to

chronic hypercortisolism. Women with CS have lower muscle mass compared to general population [192]. Decreased libido, a sign of hypoandrogenism in both genders, is reported by 40% of female CS patients[63]. But, in contrast to males, females with CS have normal or even increased AAS synthesis and levels, compared to healthy controls [193, 194]. Four fifths of women with CS have menstrual irregularities, which has been attributed to hyperandrogenism, direct cortisol action, or depletion of LH or estradiol[175]. More than 75% of CS cases present with hirsutism, that is, male-patterned body and face hair growth in female patients, and a clear sign of hyperandrogenism[1, 63]. Women with

CS-related hirsutism have androgen levels higher than healthy controls[195]. Other signs of hyperandrogenism, such as voice changes or acne, are rare in female CS.

In infants, tumors causing CS are exceedingly rare. In pediatric Cushing’s disease and adrenocortical carcinoma, AAP circulating levels are usually normal for the age[196, 197]. Virilization signs such as change in voice, penile or clitoridian overgrowth, and hirsutism are common [198]. Published studies do not describe muscle changes in these children, possibly due to difficulties in assessment.

In adult female and in pediatric CS, virilization, muscle catabolism, and circulating androgens changes are not correlated. These examples suggest that relative hyperandrogenism in some tissues may be paralleled by relative hypoandrogenism in others.For example, it may be possible that, in some tissues, excess GC activates the androgen receptor (AR)[199], the nuclear receptor specific for AAS at physiological concentrations. Because short-term Dexa inhibits AR expression in women’s muscle[200], it may be possible that GCs interfere with

Testo signals in a tissue-specific manner.

Understanding causality in the case of simultaneous muscle loss and hirsutism is complicated by dose- and compound-dependent crossconversion of GCs to AAS and interference of GCs in AAS synthesis and degradation. It is unclear to what degree muscle loss in CS is influenced by the changes in endogenous AAP and AAS. Based on endogenous levels, it appears that AAS therapy may benefit men, but not women and children, with CS.

Interestingly, the Crawford and colleagues trial observed muscle protection by

AAS as adjuvant to GC therapy although, at enrollment, these men had circulating Testo levels in the lower normal range[158]. This confirms that GC deleterious effects are not solely caused by hypoandrogenism.

Hypercortisolism is associated with hypoandrogenism solely in adult males. Androgen therapy for muscle protection in CS is predicted to benefit them more than other populations.

## Molecular mechanisms of androgenic myoprotection in humans

GAML is a phenomenon well-studied, with its molecular mechanisms dissected in human studies. In contrast, the effect of AAS in GAML was studied in a few case reports, marred by the absence of objective physical outcomes and of molecular analysis. More information can be gleaned from the effect of AAS in other muscle-depleting conditions.

Most commonly, studies of AAS on muscle are carried on men with lower than normal circulating Testo and / or associated symptoms, also called hypogonadal. In male primary hypogonadism, rates of cortisol synthesis and degradation are typically normal[201]. In this population, AAS therapy, even with low, “replacement” doses, causes an increase in muscle mass and force[202, 203]. The gain in muscle mass is caused mainly by an increase in protein synthesis, as evidenced by increased nonoxidative uptake of labeled leucine[203]. Moreover, Testo causes an increase in FSR of myosin heavy chain (MyHC), indicating that protein accretion is localized in the myotubes.

The referenced studies also measured leucine flux, a proxy for protein degradation, but failed to detect significant changes as a result to Testo therapy. The absence of a detectable change in leucine flux may be attributed to a true lack of effect on catabolism, or may be an artifact caused by the use of whole-body, rather than isolated muscle, methods.

Typical naturally-occurring male hypogonadism is usually associated with pleiotropic pathology, such as Klinefelter’s syndrome, where deficient androgen synthesis may be complicated by other peripheral defects. For this reason, some studies were conducted in males with iatrogenic hypogonadism, induced by administration of GnRH agonists, such as goserelin or leuprolide, which disrupts and eventually abolishes LH secretion. Leuprolide-induced hypoandrogenism causes loss of muscle mass in healthy volunteers and in prostate cancer patients[204, 205]. In the former, most of the muscle losses are reversed if exogenous Testo is co-administered. Chemical castration causes decreases in both protein synthesis and degradation[206], suggesting that, in some cases, such as restoration of physiological levels, Testo supplementation may be followed by a paradoxical increase in protein degradation.

The protective action of AAS therapy in iatrogenic hypoandrogenism is not affected by co-administration of an aromatase inhibitor such as anastrozole[207]. Aromatase converts Testo in estradiol. The continuing muscle protection when Testo cannot be converted to estrogens demonstrates that muscle protection is an intrinsic ability of Testo. A more plausible mediator is the anabolic hormone insulin-like growth factor I (IGF-I), whose muscle expression is decreased by iatrogenic hypogonadism[206], and by short-term, high-dose Dexa.

Another well-studies group comprises older men, whose Testo levels and muscle mass are naturally declining[208, 209]. An argument has been made about benefits of Testo replacement therapy in this population. Multiple clinical studies tested this hypothesis. In older men with low bioavailable Testo, muscle mass and strength is improved by 200 mg Testo every other week[210, 211]. As in hypogonadal men, muscle recovery can be localized to the contractile cells, as indicated by increases in the CSA of fast- and slow-twitching fibers[212]. No evidence of fiber type switching or fiber type-specific effects in response to AAS therapy has been seen. Instead, histological studies reveal that elderly treated with AAS have significantly more satellite cells[212].

Testo causes improvement in the net balance between protein synthesis and degradation at muscle level[213]. The cause of protein accretion is an increase in protein synthesis, as shown by an augmentation of mixed-muscle FSR[214]. Interestingly, some of this newly accrued protein is extracellular matrix, as indicated by the upregulation of circulating N-terminal propeptide of type III procollagen[215].

Ferrando and colleagues made the case for an anti-catabolic action of AAS in older men[3]. However, their study differs in key aspects from the other studies and the medical practice. They tested a variable, moderate dose of Testo on normogonadal older men, with the goal of maintaining a physiological Testo level. Moderate Testo therapy caused an improvement in muscle mass, strength, and net protein balance. However, they failed to observe an improvement in protein

FSR, and concluded that the net protein balance improvement must be caused by T-induced inhibition of protein degradation. The failure to detect protein synthesis rate changes indicates that, perhaps due to unusual treatments, this study yielded unusual outcomes, which cannot be extrapolated to other studies or populations.

In support of their hypothesis, Ferrando and colleagues showed a significant decrease in the rate of phenylalanine disappearance at muscle level and in the proteasomal enzymatic activity. However, the data they provide show that, on the contrary, proteasomal activity is not reduced by Testo therapy. Digitizing their plot (Fig. 1) indicates that six-months of placebo changed the normalized lactastatin-sensitive proteasome activity from 0.117 to 0.0761 relative units, whereas six months of Testo changed it from 0.0839 to 0.0784 relative units, a likely non-significant set of changes. The same group found a similar pattern of anticatabolic action, in a short-term trial of Testo on men with severe burns, once again doubled by an apparent absence of the pro-anabolic component[216]. It may be possible that the protective action of Testo changes qualitatively, depending on the cumulative dose. Alternatively, the anticatabolic action may be more salient when Testo supplementation is given to the normogonadal. The hypothesis that Testo inhibits protein degradation remains tempting, but better studies are needed.

In older men, Testo upregulated intramuscular and circulating levels of IGF-I[213, 217]. The protection of muscle force provided by Testo to the older hypogonadal men is not hindered by co-administration of finasteride, an inhibitor of 5α-reductase which causes the transformation of Testo to

5α-dihydrotestosterone (DHT)[218]. Similar lack of effect was seen with dutasteride, a less specific inhibitor of 5α-reductase, added to exogenous Testo, in a younger, possibly less hypoandrogenic cohort[219]. In human males, conversion to DHT is not required or T’s regulation of muscle mass. Possibly more relevant,

Testo upregulates IGF-I in the muscle and in the serum of the older men[213, 217].

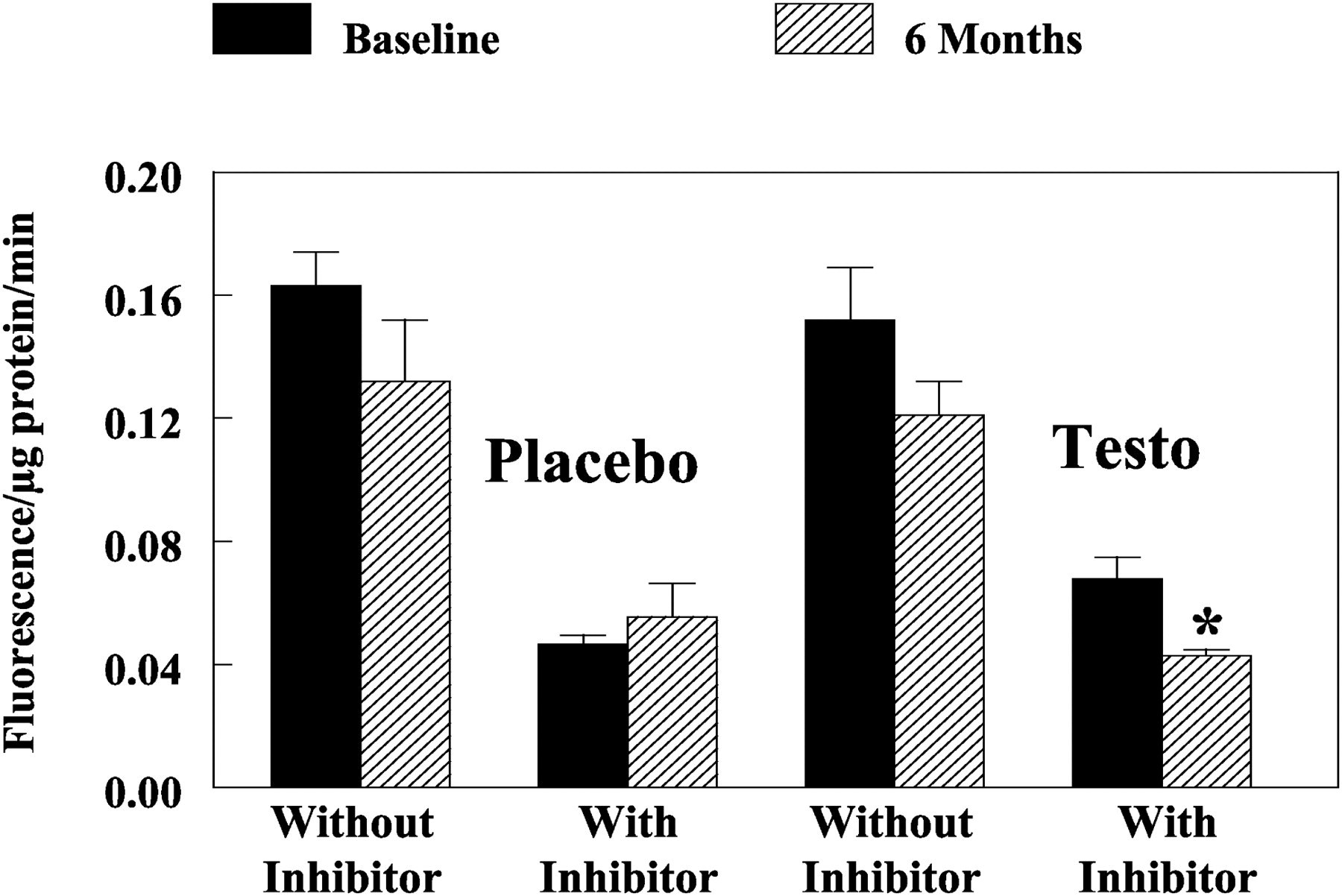


Figure 1: Changes in proteasome enzymatic activity following 6 months of Testo therapy (reproduced from [3] with permission).

While women’s endogenous AAS levels are lower than men’s, it is unclear if benefits of Testo therapy outweigh the deleterious effects[220, 221]. There is no FDA-approved Testo preparation for women. Therefore the action of Testo in women losing muscle is yet to be investigated.

The best molecular observations on the action of Testo on muscle loss have been obtained from studies of HIV-positive men, who have significantly lower circulating Testo levels[222]. Some of the drugs used in HIV-AIDS, the anticachectic megestrol and the protease inhibitor ritonavir, may cause hypercortisolism, making it even more informative about the action of AAS in CS. AAS delays loss of muscle mass in AIDS wasting syndrome, leading to better quality of life[223]. Microarray analysis indicated that T-treated muscle upregulated, as expected, expression of genes from the IGF-I- and AR-stimulated signaling pathways[224]. Immunoblot confirmatory studies indicated that Testo caused the upregulation of a key component of the IGF-I signaling pathway, the protein kinase B, also known as Akt, in its Serine-473 phosphorylated, that is, activated form. Other genes upregulated by Testo are muscle development regulators, such as the myocyte enhancer factor 2A (MEF2A) and a host of macrophage-associated markers. In addition, Testo stimulated expression of genes from other pathways, including transcription factor 4 (TCF4) from the Wnt / β-catenin pathway, AMP kinase (AMPK), and the guanine nucleotide exchange factor Sos, involved in the mitogen-activated protein kinase (MAPK) pathway. However, in the same study, at protein level, MAPK did not appear to be modulated by AAS therapy. The referenced microarray study failed to find a change in expression of the major muscle regulator myostatin[225, 226], or of the two E3 ligases typically associated with muscle loss, MAFbx and

MuRF1[227, 228].

The histological and molecular findings from hypogonadal and HIV-positive males receiving AAS have been confirmed in many other pathologies that cause loss of muscle. AAS therapy improves muscle mass and strength in males with chronic kidney disease and liver cirrhosis[229, 230]. In men with COPD, 100 mg Testo enanthate injected weekly led to improvements in muscle mass and strength, potentially augmenting quality of life[231]. These improvements are caused by an increase in fiber CSA, regardless of fiber type, and by an upregulation of the IGF-I mRNA isoform known as mechanogrowth factor or IGF-IEc[232]. In these men, MyHC upregulated by Testo in these men was of isoform 3, also known as embryonic MyHC. This is also one of the two MyHC isoforms upregulated by Testo in the HIV-positive men. In the COPD cohort, embryonic MyHC was found in thinnest fibers, possibly marking them as newly formed.

A cross-sectional study split a cohort of males with heart failure, without cachexia and with normal circulating cortisol, Testo and ACTH levels in two halves based on their cortisol level. The subgroup with lower circulating cortisol achieved a higher peak work rate, suggestive of GC-induced muscle damage[233]. Some randomized trial showed that heart failure patients improve their muscle force with AAS therapy[234, 235]. However, a series of recent studies found deleterious cardiovascular effects of AAS[236, 237], which will discourage the use of Testo in heart failure. In fact, many of the aforementioned conditions where AAS was used are marred by higher cardiovascular risks and frailty, pressing the need for alternative, equally efficient, anabolic adjuvants. To this end, a deeper understanding of AAS therapy at molecular level is required.

In various conditions that cause muscle loss, AAS benefits share a pattern including improved muscle mass and strength, fiber hypertrophy, tissue remodeling, and increased protein synthesis. In some conditions, AAS-driven muscle rehabilitation is associated with an increase in satellite cells and / or an inhibition of protein degradation. Putative molecular mediators known from animal models have not been confirmed in humans, with the invariant exception of the upregulation of IGF-I. Better clinical studies are required. Animal models and in vitro studies sketch the road ahead.

**2. BIOLOGICAL PREMISES**

## Skeletal muscle histology

Muscles are specialized for their main ability, contractility. For mammalians, ability to move is vital for survival, meaning that a large portion of their bodies is muscle. In a cohort of 300 borderline overweight US Americans, skeletal muscle as a proportion of body weight was on average 41% for men and 31% for women[238]. Owing to the ability to measure individual muscles in any experiment, the scientific community has not been under pressure to develop accurate techniques that measure total skeletal muscle mass in mice. A proxy measure of murine skeletal muscle mass, lean body mass averaged 81% of total body weight in adult males (detailed in results section). Three fifths of the human body’s protein is confined to the muscle contractile and support structures[239].

For the most part, the skeletal muscles confers the three-dimensional intricate conformation of the body, suggesting a complex, detailed organization, at least at macroscopic level. In contrast, at cell level, the relatively high specialization of the skeletal muscle leaves little space for diversity or inhomogeneity. Skeletal muscles are organized in anatomical units that may impose a force on two moving body segments (typically, bones), determining them to come closer to each other. This specificity of action is ensured by the presence of distinct, well-determined insertion points, to which skeletal muscles are attached by the means of tendons and aponeuroses (reviewed in [240]).

The tendons are dense connective tissue structure, which extend into the epimysium, a connective tissue sheath surrounding the muscle. In turn, epimysium emits connective septing structures termed perimysium, splitting muscles into subunits termed fascicles. At an even lower level, a thin, sparse

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connective structure called endomysium coats each polynucleate, elongated cell (termed myofiber). Connective tissue supports the terminal branches of the nervous, circulatory and lymphatic systems. In addition, connective tissue inside the muscle provides mechanical anchoring between fibers, longitudinally, laterally and with the tendons. This is particularly true of perimysium, whose collagen content is 95%[241]. The collagen in muscle-associated connective tissue is mainly of types I and III (reticulin), with traces of type V collagen and fibronectin, while the most frequent proteoglycan is collagen I-binding decorin[242]. The external lamina is the equivalent of basement membranes in other tissues, an proteic structure surrounding each multinucleate, roughly tubular cell. The external lamina contains collagen IV, laminin, and heparan sulfates [243, 244]. Muscle mass changes should require remodeling of all these connective structures, with novel collagen synthesis possibly misleading measurements of protein synthesis in atrophying muscle. In addition to the contractile and connective components, muscles include vascular, nervous, adipose, blood and immune cells.

The myofiber is the histological base unit of contractile tissue. Another muscle-specific population of cells comprises the mononuclear, proliferating cells, called satellite cells. Satellite cells are nearly devoid of cytosol, sitting in close proximity to the fiber, under the external lamina. They can be identified by markers such as Pax7 and, on the membrane, CD34 (reviewed in [245]).

Each myofiber is a syncytium, that forms and grows by fusion with neighboring cells. The only way to acquire nuclei to the myofibers is fusion with surrounding proliferative mononucleate cells, or with neighboring myofibers. Cells that have the ability to undergo mitosis and to fuse with myofibers are typically designated myoblasts. In vivo, myoblasts are derived from a subset of satellite cells. Experimentally, α7 integrin is an effective marker for selecting proliferative precursors from muscle[246], although its sensitivity and specificity are yet to be established. Alternate sources of nuclei in the myofiber are subject of ongoing research, but their relative importance is expected to be minor at best (reviewed in [247]). A transplant of seven satellite cells from an adult mouse is capable of yielding more than a hundred multinucleate myofibers, thus demonstrating former’s ability to regenerate muscle[248]. The transformation of quiescent satellite cells to proliferating myoblast is regulated by the interplay of growth factors, external lamina, and contact with myofibers[249]. In vivo, myofiber nuclei are typically peripheral, while in vitro incomplete models often yield elongated, contractile cells, with central nuclei, akin to regenerating fibers.

The latter are typically termed myotubes.

The proliferative niche can play an important role in muscle atrophy and recovery. However, muscle hypertrophy may occur without cell divisions. For example, the muscles of mice receiving clenbuterol and of rats undergoing eccentric training gain 20-30% muscle mass without apparent DNA changes[250, 251]. Quail muscles depleted of proliferating cells by irradiation still undergo hypertrophy in response to stretch-overload[252]. These example of amitotic hypertrophy demonstrate that, in some circumstances, the number of nuclei and the pace of transcription are not limiting factors in muscle growth.

A large majority of the myofiber cytosol is the contractile apparatus, in the shape of bundles of protein filaments termed myofibrils. Within myofibrils, myosin and actin filaments alternate, held together by multi-protein complexes containing titin. Myofibril proteins are about two thirds of the total myofiber protein[253]. Therefore, any myofiber size change with functional relevance should correlate with changes in myosin II and actin protein content. In rat muscles, three days of streptozotocin-induced acute diabetes causes intramuscular formation of a actin degradation fragment[254]. The apparent sensitivity of actin to muscle mass regulators poses a technical challenge to protein and even mRNA measurements, because traditionally actin is considered a housekeeping, unregulated, and rather constant protein, and is used in level normalization. If, in an atrophying muscle, some protein of interest is depleted faster than actin, immunoblots will convey the certainty that the former is downregulated. But if the protein of interest is lost in a less preferential manner, at the same rate with actin, immunoblots normalized to actin will convey the mistaken appearance of constancy.

Similar issues govern the use of 3MH as a marker for myofibril protein catabolism. Given that the main correlate of urinary 3MH is muscle mass[255], 3MH measurement may lack sensitivity when used as a indicator for muscle catabolic rate. On one hand, increased catabolism is expected to cause increased 3MH output, but on the other, an atrophic muscle has less 3MH to release.

During experiments that perturb muscle mass equilibrium, the level of the regulators, of typical housekeeping proteins, and of non-myofiber proteins may fluctuate in manners that convolve their specific modulation with overall muscle protein kinetics. Investigation of recovery from muscle loss is burdened by the fact that it aims to dissect protein regulation, when the regulators are proteins themselves.

## Preadult muscle development

Around puberty, muscle growth is associated with a massive shift of nuclei from the satellite cells to the myofibers[256]. Even earlier, de novo muscle development is remarkable for its accretion of new nuclei to the myofiber. Pre-adulthood muscle growth appears reliant on hyperplasia, that is, cell proliferation. Cell proliferation regulators are crucial in determination of muscle mass in before and soon after birth.

In utero, the mesoderm, which is the source of muscle progenitor cells, undergoes segmentation and differentiation to form somites, dermomyotomes, and eventually myotomes (reviewed in [257]). The latter contain the earliest cells expressing muscle regulatory factors (MRF). In animal models, the cells in epaxial and hypaxial muscle initiate the formation of trunk and limb muscles. The limb muscle early progenitors express myogenic factor (Myf) 5, due to stimulation from the transcription factor Pax3[258, 259]. Myf5 is a strong inductor of the muscle transcriptional program and phenotype, with the ability to convert embryonic fibroblasts to myosin-containing syncitia[260]. Cranial muscle formation is coordinated in a partly different manner, through the transcription factors T-box (Tbx) 1 and paired-like homeodomain (Pitx) 2 (reviewed in [261]). Once this early stage is completed, later fetal muscle progenitors converge to a phenotype remarkable for the expression of the MRF MyoD, due to stimulation by the transcription factor Pax7[262]. MyoD knockout mice are normal, with Myf5 supplanting its absence[263]. In the Online Mendelian Inheritance in Man

(OMIM) database, there is no reported case of human mutation of Myf5 or MyoD, further supporting the idea of duplicate function. Neither Myf5, nor MyoD induce muscle attributes. However, the expression of either will promote expression of another MRF, myogenin, which marks the transition form specification to differentiation. MRFs share structural features, and some are juxtaposed in the genome, making their study difficult.

Initiation of myogenin expression marks the transition from specification to differentiation stage. In cultured cells, myogenin expression is followed by p21 expression, which removes the muscle precursor from the cell cycle[264].

Subsequent changes include expression of muscle-specific enzymes and contractile proteins[265], of a fourth MRF, the myogenic regulatory factor 4 (Mrf4)[266], and, finally, acquisition of fusogenic abilities. In humans, by the seventh week of gestation, the initial wave of myoblast fusion slows down, and a second proliferative stage starts. The latter tapers off, leading to formation of secondary myotubes, within the same laminar sheath with a primary myotube. Around the seventeenth week, some secondary myotubes migrate to form independent centers of coalescence for a third set of myotubes[267].

In chick embryos, overexpression of IGF-I induces a rapid increase in the ratio of myoblast to myofiber nuclei, while fiber density is unchanged[268]. In addition to the hyperplastic effect, IGF-I stimulates protein anabolism in prenatal muscle[269]. Defects in IGF-1R signaling determine low birth weight and subsequent growth retardation in humans and transgenic mice, although it is unclear this is due to the mitotic or to the anabolic deficiency[270, 271]. Moreover, the murine model may be marred by the formation of hybrid dimers between

IGF-1R and insulin receptor, which is likely to impair insulin signaling as well.

In mice, phenotypic differences between myostatin-null and wild type develop by the second week of embryogenesis[272]. Experimental myostatin perturbations in chicken embryos alter the population of muscle precursor cells[273]. The few reports of viable human mutations in the myostatin gene concern newborns with unusual unusually large muscles, due to loss of function[274]. Nevertheless, myostatin defects appear to cause ampler changes in muscle mass after birth[275, 274]. The relative importance of myostatin in embryo muscle development is to be determined.

Immediately after birth, the number of satellite cells is much higher than in the adult, with a one magnitude order drops between birth and 10 years[276].

This decay carries on throughout the life time at slower rate. Perinatal Pax7 knockout reduces muscle ability to regenerate, while its genetic depletion in utero or at adulthood does not exhibit pathological traits[277], suggesting that juvenile muscle growth is distinct from muscle development at other ages. Overall, muscle formation in children appears qualitatively different from adult muscle hypertrophy and regeneration, and should be more susceptible to modulation via mitotic mechanisms. This may be underlie a difference between juvenile and adult in the atrophying effect of GC on muscle.

The most likely source of new myofiber nuclei are the adult satellite cells, derived from a subset of fetal Pax3+Pax7+ cells. Some cells, such as bone marrow stem cells and pericytes, have the ability to fuse with myotubes, and possibly contribute to the muscle stem cell population[278, 279]. In adulthood, most limb and trunk satellite cells are Pax7+, while head satellite cells are often Pax3+Pax7-[280]. However, in longitudinal studies, human muscle mass, as estimated by body potassium, decreases in men and and stagnates in women over the age of 30[281]. As evidenced by studies such as the New Mexico Elder Health Survey, 1993-1995, with aging, loss of muscle mass accelerates [282]. Any muscle growth mechanism, including that dependent on satellite cells, is likely less powerful in adulthood than in childhood. Indeed, after the age of 20, nuclei in human muscle maintain an almost constant length of telomeres, indicating that mitosis is a rare phenomenon in the adult muscle[283]. Moreover, the proportion of satellite cells with proliferating abilities decreases with age, as more of them approach the Hayflick limit[284].

## Physiological muscle metabolism

Muscle is a major energy user in the body, using fat during rest and glucose during exercise (reviewed in [285, 286]). Because its capacity to synthesize fatty acid is negligible, muscle is a consumer and a minor store, but not a generator, of fatty acids. During fast, more than half of infused non-esterified fatty acids (NEFA) are taken up by muscle, with a higher rate of incorporation in type I oxidative muscle[287]. In contrast, after feeding, oversupply of NEFA is compensated mainly by increased uptake, in absolute terms, in the visceral adipose tissue, with minor contributions from other adipose tissue, liver, and muscle. After repeated exercise, muscle lipoprotein lipase expression is increased, indicative of an adaptive improvement in muscle ability to extract NEFA from circulating triglycerides[288]. Ongoing studies suggest that muscle oxidative (catabolic) uptake of NEFA is upregulated by peroxisome proliferator-activated receptor (PPAR) β/δ, which stimulates expression of the lipolysis rate-limiting enzyme, carnitine palmitoyltransferase I[289, 290]. This hypothesis is supported by fast- and exercise-induced upregulation of muscle PPAR δ[291, 292]. Moreover, PPAR δ overexpression leads to increase in type I fibers and subsequent resistance to high-fat diet[289]. Organ-level studies are impaired by the existence of nontrivial intramuscular adipose tissue.

This work uses extensively the C2C12 cell line, an immortalized female mouse muscle progenitor line obtained from a muscle recovering after mechanical injury. Treatment of confluent C2C12 cells with Dexa and 3-isobutyl-1-methylxanthine (IBMX) causes their differentiation into adipocytes[293]. Some pre-adipocyte traits, such as upregulation of PPAR γ, Krüppel-like factor-15 (KLF-15), and CCAAT-enhancer-binding protein (C/EBP) β and δ ([294]; reviewed in [295]), may surface in cell culture experiments where muscle differentiation did not complete, and residual potential for adipogenesis remains (see, for example, [296]). An in vitro shift to a more adipose-like phenotype may be associated with diminished cell fusion ability, lower protein synthesis and lower mitochondrial content, which may be misinterpreted as muscle atrophy.

In humans, up to 90% of the glucose absorbed after a meal is removed from circulation by the skeletal muscles[297, 298], meaning that muscles should have a paramount role in the development of insulin resistance and eventually diabetes mellitus. After normal feeding, muscle builds polysaccharides reserves, in part because it can synthesize and deposit the largest glycogen stores in the body, but also because it cannot release glucose.

Muscle work is generated at such high rates, that most glucose is processed solely through glycolysis, in the cytosol, to the three-carbon pyruvate. Some of the pyruvate is further oxidized in the muscle, through the tricarboxylic acid cycle, but a significant amount is converted to lactate and released in the blood stream. As part of the Cori cycle, circulating lactate is reassembled into glucose by the liver, and re-released into the blood stream, for muscle use. A similar shuttling mechanism further enables muscle to rely on glycolysis, by transaminating excess pyruvate to the amino acid alanine, similarly released in the blood. Circulating alanine is converted by liver to glucose. Pyruvate transamination requires the amino acid glutamate. Muscle uses glutamate for other metabolic processes, including the synthesis of non-essential amino acids, including proline and arginine. Therefore, at rest, human muscle uptakes significant amounts of glutamate, less serine, while releasing alanine, glutamine, and smaller amounts of the other amino acids[299]. Glutamine is synthesized in muscle in order to release the excess nitrogen yielded by amino acid release during protein degradation.

Muscle’s glutamine is then converted by the liver to urea and excreted.

Among the amino acids with a trend for release between meals, isoleucine, leucine, methionine, phenylalanine, threonine, and valine cannot be synthesized by humans. Their net release indicates that, at rest, basal level of protein degradation slightly surpass protein synthesis. Conversely, for a steady muscle mass, there must be net protein synthesis in the fed state. Feeding status is relayed to the muscle by a surge in insulin and, independently, by an increase in circulating levels of essential branched-chain amino acids (BCAA), that is, valine, isoleucine, and leucine. The signaling effects of BCAA are not completely understood. Some studies show that BCAA ingestion or infusion increase protein synthesis rate[300], while others claim that BCAA solely reduce protein degradation[301]. However, staying true to their energetic value, BCAA supplementation caused increases in insulin in all the referenced studies. It becomes difficult to extricate BCAA intrinsic effect from that of insulin. Moreover, the combination of insulin and BCAA is a potent synergistic anabolic stimulus in healthy human muscle[302]. One study investigated the action of BCAA at clamped normal insulin levels[303]. In young, healthy controls, BCAA alone were able to increase the fractional synthesis rate for myofibrilar protein. At the same time, BCAA caused an increase in phosphorylation of mTOR at Serine 2448. This posttranslational modification is caused by the ribosomal protein S6 kinase, 70 kDa (p70-S6K)[304]. Interestingly, p70-S6K is a substrate of mTOR complex 1 (mTORC1), with the latter considered an integrator of nutrients, energy, and growth factor signaling (reviewed in [305]). Indeed, in the same muscle, p70-S6K was activated, as indicated by an increase in its Thr 389 phosphorylated form. Another substrate of mTORC1, 4E-BP, was hyperphosphorylated. The action of mTORC1 on 4E-BP is the canonical way in which the former stimulates protein synthesis, by abolishing the latter’s ability to bind and inhibit the mandatory translation initiation factor eIF4E. Many of these anabolic markers may be paradoxically elevated in acute, non-starving atrophy settings, where BCAA sudden release increases their circulating levels.

In addition to their signaling roles, BCAA are also used by muscle as protein precursors, and as energetic substrates, when preferred energetic substrates are not available[306]. In studies which measure leucine disappearance from the bloodstream or culture medium, a distinction must be made between the leucine used in non-oxidative, anabolic reactions and the alpha-ketoisocaproate-forming, ergogenic usage.

The degree to which energetic needs modulate BCAA usage appears to differ between humans and rats, reducing the validity of animal models (reviewed in [307]). Amino acid uptake or release rates are perturbed by energy and hormonal factors, to a degree that is still to be measured in humans. An argument has been made for using phenylalanine as a tracer, because muscle catabolism is negligible, and because it has a lower insulin secretagogue effect[308, 309].

## Adult muscle remodeling

With aging, muscle gradually shifts from a mitotic to a postmitotic profile, with muscle growth achieved increasingly through hypertrophy, that is, cell size growth. While the typical middle-aged or elderly adult is undergoing net loss of muscle mass, the muscle is still adept at re-growing and remodeling under two common circumstances, exercise and injury.

Humans achieve muscle mass growth following exercise, although some forms of exercise are more suitable at increasing strength or resistance that mass per se. In animals, muscle growth is induced by muscle overload, or muscle unloading and reloading, which may be conceived as forms of aerobic exercise. In healthy volunteers, the acute response to exercise includes increased intramuscular expression of MRFs MyoD and myogenin, and increased circulating IGF-I and IL-6[310, 311, 312]. These signals are associated with increased proliferation of satellite cells and recruitment of neutrophils to the muscle[313, 314]. In the acute phase, the satellite cells colocalize with IGF-I[310]. The negative muscle regulator myostatin is not correlating with the phenotype, that is, it is not decreased by acute exercise[314, 315]. Acute exercise increases the fractional protein synthesis rate in muscle[316, 317]. Interestingly, a single bout of exercise during fast leads to increases circulating cortisol levels and the release of

3-MH, indicative of increased stimulation of protein degradation[318]. Concomitant amino acid feeding extinguishes GC and catabolic response to exercise. Variations in regimens of exercise and timing and composition of diet led to a plethora of studies. Just as most of the short-term exercise routines do not lead to muscle macroscopic changes, some studies describe an absence of changes at molecular level following short-term exercise (reviewed in [319]). An important future direction in exercise science is establishing what distinguishes an effective brief exercise routine from an ineffective one. In this context, GC may be interesting noninvasive markers.

In the long term, exercise increases fiber CSA, density of satellite cells, and the number of myofiber nuclei, while the level of intramuscular MyoD, IGF-I, myostatin slowly return to normal[320]. On the other hand, long-term exercise induces the expression of catabolic markers, such as the E3 ligases MAFbx and MuRF1[321]. Taken together, these molecular findings indicate exercise causes muscle remodeling, which manifests as increased muscle turnover, with upregulation of both protein degradation and synthesis. Moreover, post-exercise muscle accretion combines hyperplasia and hypertrophy.

A similar biphasic response is yielded by injury. In the immediate stage after injury, the muscle is infiltrated by pro-inflammatory M1 macrophages, while at later stage, anti-inflammatory (M2) subclass dominates (reviewed in [322]). Although the studies are rather incomplete, it appears that, similar to exercise adaptations, injury triggers a burst of growth factors, probably including IGF-I, basic fibroblast growth factor (bFGF), and transforming growth factor-beta (TGF-β) ([323]). Most of the studies of regeneration provide circumstantial evidence, such as improved healing in the presence of a presumed mediator, rather than impaired healing in its absence. Still unidentified molecules from crushed muscle are able to cause myoblast hyperplasia, above the levels caused by stimulation with known growth factors[324].

Less than half of the C2C12 cells in their proliferating, undifferentiated form, express MyoD or Pax7[325]. Limited evidence suggest these myoblast-like cells do not express Pax3 either[326]. Therefore, the C2C12 line are an incomplete model of hyperplastic muscle accretion.

Generation and regeneration of muscle in common scenarios, such as development and adaptation, remain an object of study, due to their complexity. Variable importance of the immune cells, of MRFs and of IGF-I, and concurring redundancy forestall attempts to envisage a common pattern of muscle hypertrophy.

## Hormonal control of muscle mass

The variability of muscle mass within population is reflective of the variable needs for muscle strength. Muscle mass and strength are adjusted to the needs of the organism mainly through humoral mechanisms. The most relevant endocrine regulators of muscle mass, AAS, GC, and IGF-I, will be discussed in dedicated sections.

While anabolic role of IGF-I at muscle level is undisputed, insulin’s effect is inconsistent, because it is easily altered by external factors. The acute phase of insulin response poses a conundrum, with some studies showing it stimulates protein synthesis in human muscle[327], while others demonstrating that its effect is limited to anti-catabolism[328]. In male rat muscle, 30 minutes in 30 nM insulin or IGF-I are equally able to stimulate protein synthesis and to inhibit protein degradation[329]. Both effects are above 10%, although the effect on protein degradation appears less ample and is marred by higher variability.

Insulin and IGF-I pathways overlap to a some degree. The liver is the main source of circulating IGF-I, under the pituitary stimulation with growth hormone (GH). However, auto- and paracrine secretions may full supplant the absence of hepatic IGF-I in adult conditional knockout mice[330]. In contrast, insulin is secreted solely by the pancreas. Our understanding of the regulation of insulin secretion is improving, dispelling the simplistic view that nutrients alone are its sole modulators (reviewed in [331]). Therefore, both IGF-I and insulin emerge as anabolic stimuli, responding to increased demand for macromolecules and / or increased supply of nutrients, with IGF-I embracing a more localized and insulin a systemic, integrative role. For both hormones, physiological concentrations are tens of times higher than the half maximal effective concentration (EC50) for their receptor, suggesting that physiological fluctuations cause marginal effects downstream[332, 333]. On the other hand, insulin has the ability to bind and activate IGF-I receptor (IGF-1R), with an EC50 about an order of magnitude lower than physiological insulinemia. The converse is true, with IGF-I being able to bind and activate insulin receptor (IR), isoforms A and B. There is a small, but real, potential for interference between insulin and IGF-I signals.

The levels of bioavailable IGF-I are under complex regulation (reviewed in [334, 335]). IGF-I may be sequestered by IGF-I binding proteins (IGFBP), which are secreted by muscle under IGF-I stimulation ([336]). The interaction with IGFBP may prevents IGF-I from interacting with receptor, or it may extend its circulating half-life by protecting it from degradation. Depending on the isoform and location of IGFBP, the interaction may result in extinction or amplification of the IGF-I signal. IGFBPs levels are modulated by insulin and their affinity is modified through competition by insulin-like growth factor 2 (IGF2). The latter can also stimulate IGF-1R, thus providing its own anabolic and pro-myogenic signals[337]. IGF2 is irreplaceable in fetal development[338], suggesting there might be distinct, unidentified receptors for this hormone family.

There is no consensus with regards to the ability of GH to stimulate muscular secretion of IGF-I. Multiple studies found an upregulation of its mRNA[339, 340], but protein data are lacking. Medium conditioned by GH-stimulated C2C12 cells fails to elicit hypertrophy in other C2C12 myotubes[341], suggesting that, at best, IGF-I has an intracellular autocrine action. In addition to the indirect effect mediated by hepatic and the putative muscular IGF-I, GH has an IGF-I-independent effect on muscle. For example, knockout of GH receptor impairs body growth further beyond IGF-1R knockout[342]. In the context of pituitary pathology associated with Cushing’s disease, the associated GH perturbations may contribute to loss of muscle.

Hypothyroidism is often associated with muscle weakness and pseudohyertrophy[343]. Other hormones such as the parathormone, are likely to have small effects on muscle protein metabolism, essentially irrelevant outside their respective pathologies[344]. In conclusion, muscle mass homeostasis is under a tight, multifactorial hormonal control, whose study is complicated by significant redundancy. The absence of third-party organs, such as glands, from reductionist cell-culture may limit their ability to replicate in vivo phenomena.

## Interaction of muscle mass and vascularization

Muscle vascularization is a modulator of muscle mass and contractility. Mice whose muscle VEGF-A secretion was genetically depleted still express a tenth of the muscle VEGF-A protein, but have only half of the capillaries per muscle fiber,

compared to their Cre-/- siblings[345]. The muscle-restricted VEGF-A-depleted mice have 12% lighter gastrocnemii, although the muscle loss disappears when muscle mass is normalized to total body weight. The loss does not affect a specific fiber type more than the others. Therefore, the muscle depleted of VEGF-A is less able of endurance effort (80% shorter time to exhaustion on the inclined treadmill) and of brief anaerobic exercise (34% lower maximal running speed).

Conversely, murine muscles injected with VEGF-A-expressing retroviruses display a higher proportion of hypertrophic fibers than those expressing bacterial β-galactosidase[346]. Moreover, in the VEGF-A-overexpressing muscle, many of the fibers have central nuclei, a sign of increased fusion with myoblasts. In the murine C2C12 cell line, VEGF-A causes faster differentiation, into longer myotubes, with more nuclei per fiber, although mitotic rates are in fact diminished. These observations are consistent with direct anti-apoptotic and profusogenic effects.

It has been suggested that the most direct effect of VEGF-A depletion is partial segregation of muscle from blood-carried endocrine signals. For example, VEGF-A-depleted muscle has lower glucose uptake, a defect which is reversed by explantation [347]. In addition, vascularization defects may induce relative intramuscular hypoxia, which is an independent atrophying, pro-proteolytic factor[348].

The relative importance of VEGF overexpression during myogenesis is still open to debate, as muscle-restricted VEGF receptor (VEGFR) knockout animals were not studied yet. Multiple effects concur to obfuscate VEGF action in hypertrophying muscle. First, VEGF promoter contains three binding sites for MyoD, meaning that growing muscle will express more VEGF[349]. Development of vasculature in growing muscle may be a physiologically meaningful way to ensure vasculature remains competent upon increase circulatory demands.

Second, although the canonical upregulator of VEGF is hypoxia-induced factor 1α

(HIF 1a), muscle VEGF is also stimulated by PPAR γ coactivator 1

(PGC-1)[350, 351]. Even in the absence of VEGF, PGC-1 coactivators facilitate mitochondria biosynthesis, leading to oxidative fiber hypertrophy and improvements in endurance capacity [352]. Therefore, it is difficult to distinguish VEGF-induced muscle changes from the common muscle remodeling program. Finally, VEGFR activation has multiple effects, including phosphorylation of Src family proteins[353], of phospholipase C (PLC) γ[354], and, indirectly, of regulatory subunits of PI3K[355] and of the Signal Transducers and Activators of Transcription (STAT) STAT3 and STAT5[356]. Most of these VEGF effects overlap with the effects of many other muscle anabolic agents. Thus, it is unclear whether VEGF plays a mediating role in muscle hypertrophy, or it provides the homeostasis of the vasculature to fiber ratio.

## Control of muscle mass through innervation

Progress made in the last few years distinguish denervation atrophy as one of the best studied models of muscle atrophy. Current understanding and future directions for the study of GAML are guided, to a large extend, by the data obtained in denervation experiments.

Limb and trunk muscles are controlled by the lower (alpha) motoneuron, located in the ventral horn of the spinal cord. Their main point of contact is a chemical synapse, the neuro-muscular junction (NMJ). In addition to the direct synaptic activity, muscles and lower motoneurons are involved in a mutually beneficial life-long interaction. The loss of NMJ-mediated stimulation leads to rapid, sizeable loss of muscle mass, commonly seen in spinal cord injury. Experimentally, denervation of a hindlimb or hemidiaphragm is a relatively simple procedure, allowing the use of an animal as its own control. Experimental denervation is the most studied model of muscle atrophy, generating working hypotheses for the study of GAML.

In utero, experimental destruction of motoneurons with bungarotoxin abolishes formation of secondary myotubes, although it has limited effect on the formation of primary fibers[357]. Conversely, motoneuron precursors from the embryo spinal cord degenerate and die if they cannot engage in significant interactions with myofibers[358]. Eliminated motoneurons include neurons whose axons fail to reach myotubes and neurons which eventually fail to maintain contact with myotubes, due to synapse elimination. The latter is a selective, competitive process, whose molecular basis is still unknown, and which ensures injectivity of the correspondence between lower motoneurons and myofibers (reviewed in [359]). For a few days after birth, rat motoneurons go into a particularly sensitive state, when axotomy determines motoneuron death[360]. Thence, axotomy elicits the reprogramming of the neuron into a less differentiated state, followed by axonal regrowth (reviewed in [361, 362]). The re-establishment of contact between the motoneuron and muscle causes the reverse molecular changes, suggesting the existence of muscle-secreted neurotrophic factors. The ability of NMJ to regenerate or even preserve its optimal structure degrades with aging, although it is not clear whether NMJ cause or are caused by aging-related muscle loss[363]. The improvement in NMJ regeneration through exercise has been seen as an important path towards its understanding. Factors that improve muscle’s ability to exercise or even mimic exercise, such as IGF-I or androgens, have been shown to benefit NMJ recovery or to slow down its degradation during aging[364, 365, 366].

The loss of contact with the lower motoneuron elicits similar changes in the muscle, with clinical relevance in the understanding and treatment of spinal cord injury and spinal muscular atrophy. Conceptually, denervation is distinct from disuse, such as that induced by damage to the upper motoneuron. Clinically, the latter manifests differently, through a syndrome termed pyramidal weakness, that mainly affect muscles opposing gravity[367]. In humans, after upper motoneuron damage, disuse response occurs in a few days after injury, and leads to exaggerated spasticity[368]. Understandably, literature does not describe any animal model of disuse by experimental damage to the upper motoneuron. In fact, literature contains multiple examples where “disuse” is taken to mean “absence of action potentials, due to denervation”, “lack of work, due to denervation, unloading, immobilization”, or even both[369]. Consequently, there are no experimental attempts to disentangle muscle-maintaining effects of work, of the NMJ transmitter, acetylcholine, and of any other musculotrophic neuron-released factor.

Experimental denervation causes rapid and ample loss of muscle mass. For example, three weeks after sciatic nerve removal, the tibialis anterior muscle halves in weight[370]. Fewer than one in 7,000 myofiber nuclei undergo apoptosis in this time[371], indicating that denervation and the fast recovery that follows renervation are more similar to the adult exercise remodeling paradigm, that to the proliferation-dependent regulation of the infantile muscle. Supporting the hypothesis on NMJ-independent trophic factors, denervation did cause massive and rapid apoptosis among stromal cells.

In a seminal study, Goldberg demonstrated that denervation leads to increased loss of prelabeled muscle protein, proving that denervation upregulates protein degradation[372]. Moreover, denervation increases urinary 3MH, indicating activation of myofibril catabolism[373]. The rate of muscle loss in denervated muscle is halved by the proteasome inhibitor bortezomib[374]. Denervation stimulates persistently proteasome enzymatic activity[375], and upregulates all the ubiquitin-proteasome pathway components, including ubiquitin, the E3 ligases MuRF1 and MAbx, and the proteasomal subunit A1 [376, 227]. The upregulation of the E3 ligases is induced by multiple independent transcription factors, including myogenin and the Foxo class[377, 378, 379, 380].

In the first few days of denervation, myogenin is induced by histone deacetylase (HDAC) 4[381]. After the first week, Foxo activation is attributed to the downregulation of its negative regulator, Akt[379, 382].

Because the mTOR inhibitor rapamycin prevents fiber hypertrophy that normally follows in vivo injections with a plasmid coding constitutively active Akt[382], the scientific community assumed, since the beginning of century, that denervation-induced Akt inactivation leads to loss of downstream mTOR-mediated effects (for example, [383, 384]). Recent studies contradicted this paradigm. In 2013, Quy and colleagues found that denervation increased Thr 389 phosphorylation and catalytic activity of p70-S6K, indicating that denervation causes in fact activation of mTORC1[385]. In 2014, Tang and colleagues proved that rapamycin, an inhibitor of mTORC1 which lacks intrinsic anabolic properties, abolishes denervation-induced loss of muscle mass[379]. These experiments prove that mTORC1 should be activated for denervation-associated muscle atrophy to proceed in its later stage. Tang showed that denervation causes phosphorylation of insulin receptor substrate (IRS) 1, proving that, in denervation, activated mTORC1 inhibits Akt through a p70-S6K IRS 1-mediated negative feedback loop. Quy found that mTORC1 activation is lost upon proteasome inhibition with bortezomib. One can hypothesize the existence of an mTOR negative regulator which is specifically targeted by the ubiquitin-proteasome system during denervation. However, the late timing of the Akt / mTOR involvement suggests that the most plausible explanation places mTOR activation after the proteasome releases significant amounts of free amino acids. It is therefore more likely that denervation-induced mTOR activation is caused by the proteasome-stimulated amino acid release.

Establishing the role for mTORC1 in denervation-induced loss of muscle mass is crucial. In conceptual frame before Quy and Tang experiments, mTORC1 was thought as being inactivated by denervation, thus downregulating on protein synthesis and removing a restriction on autophagy. This hypothesis was conceptually attractive, because it implied a co-operation between multiple anabolic and catabolic pathways, towards achieving muscle loss. In particular, denervation-induced autophagy appealed to the muscle biologists of the 2000’s. For example, in a never-replicated experiment, one group claimed that the lysosomal inhibitor chloroquine prevents denervation-induced muscle loss[386]. Others showed that denervation upregulates lysosomal enzyme cathepsin L[387], while a third group claimed that denervation causes buildup of the autophagosomal marker, microtubule-associated protein 1 light chain 3 (LC3)

[388]. However, most experiments refuted the role of autophagy in GAML. The lysosomal inhibitors leupeptin, methylamine, and E64-c have minimal effects on the release of free tyrosine from denervated muscle[373]. Transgenic mice expressing LC3 fused with green fluorescent protein (GFP) exhibit a loss of autophagosomes in denervated muscles[385].

A similar debate surrounds protein synthesis regulation in denervation. In Goldberg’s 1969 experiment, the specific activity of the remaining muscle protein was essentially the same as in control limbs. Increased or even unchanged protein synthesis rates would have caused a reduction in specific activity. Therefore, Goldberg concluded that denervation causes protein synthesis decreases, thus opening a debate that is still unsettled. Goldberg’s observed reduction in protein synthesis rate is consistent with an indiscriminate loss of translation initiation and elongation factors, as much as with an active mechanism of protein synthesis cessation. Moreover, more recent functional studies, including some from Goldberg’s group, found that denervation stimulates translation[389, 385, 375, 390, 373]. Unlike the 1969 study, contemporary studies measured protein synthesis rate directly and over shorter time intervals (hours, rather than weeks). As in the case of autophagy, follow-up studies found molecular changes in regulators of translation, in some cases supporting a regulated, denervation-triggered mechanism of protein translation shutoff, in others refuting them. Both Tang and Quy found that denervation causes increased phosphorylation of 4E-BP, thus leading to stimulation of protein synthesis. Both also show that the effect on 4E-BP is prevented by rapamycin, demonstrating that protein synthesis is upregulated through an mTOR-dependent mechanism.

Given that rapamycin has essentially no effect on adult muscle mass, it may be that mTOR is not the crucial effector of denervation-induced muscle loss once thought. Recent studies show that its activation, and the downstream net anabolic effects it elicits through autophagy and translation, cannot explain the observed phenotype. Denervation-induced muscle loss is, to great extent, produced by the ubiquitin-proteasome system. Better experiments are needed in order to understand autophagy and of mTOR in denervation-induced muscle loss. The irreconcilable differences between functional and molecular studies of denervation epitomize similar predicaments in the study of muscle atrophy due to other etiologies, including the less-explored GAML.

## Animal models of glucocorticoid myopathy

In the 75 years since the discovery of chronic steroid myopathy, scientists attempted, with variable success, to develop multiple animal models. In dogs, seven days of 0*.*44 mg/(kg d) cause increased nitrogen excretion, increased glutamine and alanine release from the hindlimb[391]. In horses, the action of Dexa on glycogen regulating pathways was replicated, but muscle mass was not measured[392].

A few species are particularly unsuitable for modeling GC myopathy. In cows, GAML is undetectable at macroscopic level. Any putative change in myofibers is compensated by bovines’ unusually rich intramuscular adipose component, further hypertrophied by Dexa[393]. In contrast to rodents, a putative mediator of GAML, myostatin, is downregulated in bulls’ muscle by Dexa[394]. In rabbits, four days of 1 mg/(kg d) did not change muscle mass and reduced urinary 3MH, despite the upregulation of some catabolic markers[395]. Another article describes GAML in rabbits, and attributes it to, unusually, lysosome upregulation[396].

For large animals, cats, or dogs, practical and humane reasons prevailed, meaning that data about muscle mass changes are often lacking. For many of these species, the genome was not known, and specific antibodies are not manufactured, meaning that signaling pathways could not be dissected. Especially in the early years, studies did not record most relevant outcomes, such as muscle mass and / or force, putting into question their validity. In the era of genome sequencing, studies of GAML focused on mice and rats. Muscle mass in mice is smaller, making dissection harder and changes closer to detection threshold, compared to rats. In terms of glucose metabolism, mice are more GC-resistant than rats[397]. It is unclear whether this resistance extends to protein metabolism. The first study of mouse GAML was published in 1964[398]. Nevertheless, rats were preferred to mice for modeling steroid myopathy. In the last two decades, mice studies became more interesting, as genome manipulations were more easily induced in mouse than in rat.

Myofiber-restricted knockout of GR abolishes GAML, while having no effect on denervation atrophy[399]. In rat muscle, chronic Dexa treatment upregulates expression of the NMJ essential component, muscle-specific nicotinic acetylcholine receptor, and resistance to NMJ-specific non-depolarizing muscle relaxants[400]. If anything, GAML appears associated with an improved NMJ, suggesting that GAML is not mediated by the motoneuron. Similarly, there is no evidence of satellite cell dysfunction in GAML. In one report, Dong and colleagues hypothesize a “glucocorticoid-induced satellite cell dysfunction”, but describe solely that Dexa reduces the number of satellite cells in a post-injury regenerating muscle[401]. In conclusion, GAML is the direct effect of Dexa on myofibers.

The perception that myotubes response to Dexa is a complete model of GAML inspired many reductionist in vitro models. Many published studies documents the effect of Dexa on the mouse cell line C2C12 and the rat cell line L6. However, these and other myogenic cell lines have significant limitations, which may cause divergence between in vitro models and the steroid myopathy they aim to describe. As mentioned earlier, in the case of primary cells, experiments with

Dexa are outright impossible, because primary cell survival is GC-dependent.

Moreover, Dexa has hyperplastic and hypertrophic effects on myogenic cell lines. Commonly tested doses of Dexa, in the range of tens of nM, have been shown to synergize with IGF-I [402] and even to act directly towards improved proliferation of L6 myoblasts[403, 404]. C2C12 fusion is more efficient when Dexa is added to IGF-I[91].

Until the end of the 1990s, Dexa was a common ingredient in myogenic culture media[405]. Even contemporary standard proliferation media, containing 10% fetal bovine serum, provide significant and unpredictable concentrations of GC, insulin, and IGF-I, impairing their study at physiological concentrations. In fusing C2C12 cells, IGF-I and Dexa synergize to amplify some Dexa catabolic signals, such as the expression of REgulated in Development and DNA damage responses-1 (REDD1)[91]. Conversely, the synergy with Dexa amplifies some of insulin and IGF-I effects, such as Akt phosphorylation on Ser 473[405]. In addition to myotubes, C2C12 may differentiate into adipose or osteoblast phenotypes. Whenever possible, this section will refer to in vivo studies. Evidence from in vitro studies on myogenic cell lines will be limited to studies where multi-nucleate myotubes were obtained and myoblasts were depleted.

With this caveat, in vitro studies have been vital in excluding third-party organs, such as the pancreas, from the analysis of GAML, at the time when genome manipulations were not available. Some molecular features of GAML have been initially described in cell culture, and confirmed in vivo later. Fully differentiated myotubes from L6 and C2C12 cell lines lose more than a quarter of their diameter when treated with 100 nM [406].

The effects of 7-day, 5 mg/(kg d) treatment on rats allow the classification of GCs in two subsets[407]. Members of the short-acting subset, including prednisolone and corticosterone, cause net gains in body weight, through increased adiposity, MR-mediated water retention, and a net neutral effect on muscle mass. In contrast, long-acting, specific GCs, such as Dexa, betamethasone, and triamcinolone, cause loss of body weight, and an even more rapid loss of muscle mass. The muscle mass loss is paralleled by reductions in maximal twitch and tetanic force[408]. GAML is an organized process, lacking the microscopic features of necrosis[409]. Although Dexa temporarily reduces food intake, possibly through stimulation of leptin secretion[410], pair-feeding experiments demonstrated that the GAML is not solely the effect of changes in appetite[411, 412]. In response to Dexa, rat myofibers undergo reductions in CSA, to an ampler degree in fast twitch fibers[413, 414]. Dexa-induced loss of muscle mass is present, although less manifest, in female rats, possibly because, in males, hypercortisolism is compounded by a reduction in circulating Testo levels[413].

The nitrogen imbalance induced by Dexa slows down by the third day, and is compensated around the seventh day of treatment[415, 416]. Given that later time points are marred by feedback mechanisms and by animal mortality, almost no published experiment on rats extends past two weeks. For an animal weighing 300 times less and aging 20 times faster than humans, the common 5-10 days experiments are comparable to chronic exposure in humans. Rats appear more resistant than humans to GAML, given that reported experiments start at about

0*.*5 mg/(kg d) Dexa, more than twice the muscle-imparing dose in humans. Illustrating the higher resistance to Dexa, published mouse experiments used higher doses (3 mg/(kg d) to 10 mg/(kg d)[417, 418]).

The next sections are informed by experiments with chronic (5-10 days) Dexa

on adult (not aged) male rodents, which are the best model from human male steroid myopathy. The few exceptions will be identified as such.

## Glucocorticoid stimulation of ubiquitin-proteasome system

The loss of muscle in the Dexa-treated adult rat is the result of an increase in protein degradation. The 1969 Goldberg study on atrophy revealed that cortisone induced an even ampler upregulation of proteolysis than denervation[372]. His comparison of fast-twitch plantaris and slow-twitch soleus found that the latter did not exhibit lower muscle mass, not increased proteolysis. Goldberg found that cortisone increases in equal manner the degradation of older and newer proteins, and of myofibril and sarcoplasmic proteins. A search begun for an undiscriminating proteolytic machine stimulated by GC.

In rats receiving 0*.*5 mg/(kg d) Dexa for six days, epitrochlearis muscle proteolysis rate increased by 50%, while protein synthesis was essentially unchanged[411]. Dexa-stimulated proteolysis affects the contractile apparatus, as indicated by the doubling of urinary 3-MH output[419]. The upregulation of proteolysis occurs even when explants, rather than animals, are treated with

Dexa, suggesting that GAML does not require extramuscular inputs[420].

In vivo GAML is best correlated with an upregulation of the ubiquitin-proteasome system. The increase in proteolytic rate is unchanged when explants are treated with the lysosome inhibitor methylamine and the lysosome / calpain inhibitor E-64. On the other hand, the Dexa-stimulated increase in proteolysis is abrogated when the explant is treated the proteasome inhibitor MG132[421], or depleted of ATP by a combination of the mitochondrial decoupling agent dinitrophenol and the unlysable glucose homologue,

2-deoxyglucose[422]. Proteasome chymotrypsin-like catalytic activity is doubled by Dexa[423]. Dexa upregulates proteasomal subunits such as C1, C2, C4, C5

[411, 421].

In cultured myotubes treated with 100 nM to 1000 nM D, the loss of diameter is reliably underlined by an 20% increase in protein degradation rate[424, 406]. Between 78% and 95% of the Dexa-induced proteolysis augmentation is lost, when proteasome inhibitors, such as beta-lactone or MG-132, are co-administered[424, 425]. Demonstrating proteasome’s primacy, MG-132 has this overriding effect even co-administered as an addition to a cocktail of lysosome and calpain inhibitors[421]. Co-administration of dinitrophenol essentially abolishes Dexa-induced proteolysis[424]. GAML dependence on ATP reinforces the idea that GC-stimulated proteolysis takes place in the proteasome. In contrast, lysosomal inhibitors had no effect, while E-64 has minimal effect[424].

In L6 myotubes, Dexa causes increased expression of the ubiquitin gene UbC, through a putative SV40 promoter-specific 1 (Sp1) response element[426]. In vivo, upregulation of UbC was confirmed only for acute Dexa treatment[420] and in diabetic rats[427].

One of the most tempting hypotheses attributed a role to NF-κB, a mediator of cancer cachexia (reviewed in [428]). However, the few reports are contradictory. In L6 myotubes, Dexa stimulates acetylation and nuclear translocation of the translational activator p65[429], seemingly in consonance with cancer cachexia. But in contradiction to such parallels, Dexa was shown to inhibit NF-κB signals by upregulating inhibitory κBα (IκBα)[430]. Moreover, Dexa’s inhibition of NF-κB was shown to be necessary for proteasomal subunit C3 upregulation in the same L6 cells[431].

Unbiased searches in atrophying mouse muscles revealed two upregulated genes, termed atrogenes, MAFbx and MuRF1[227]. Both are E3 ligases, pointing to an important role for the proteasome-ubiquitin system in muscle atrophy.

In contrast to other muscle atrophy models, GAML is associated with stronger reliance on MuRF1 than on MAFbx. While genetic depletion of either atrogene leads to muscle sparing in the denervation model[227], only MuRF1, but not MAFbx genetic depletion spares muscle treated with Dexa[432]. The incomplete sparing of the MuRF1 knockout may be caused by genetic redundancy, but may also be explained by a need for a second, co-operating gene.

While MuRF1 promoter includes a GC response element, no similar structure was found on MAFbx promoter[433, 434]. Indicating a lower amplitude and / or higher variability in MAFbx, unbiased searches in rat GAML failed to identify MAFbx as a target of Dexa[435].

The difference in amplitude of change may be related to the role that MAFbx plays in muscle restructuration. In C2C12 cells, MAFbx is upregulated by differentiation[436]. MAFbx is induced in muscle during hypertrophy from reloading[437]. Moreover, MAFbx knockout abolishes hypertrophy of loading[438]. In contrast to denervation atrophy, it appears that remodeling in GAML is minimal. Supporting this distinction, denervation is a stronger upregulator myogenin than Dexa[439]. Since MAFbx promoter is activated by myogenin [378], the moderate myogenin response in GAML may explain a less ample MAFbx upregulation.

Substrate specificity is also distinguishing the two ligases. The only two known MAFbx ubiquitination substrates are MyoD and the eukaryotic initiation factor 3f (eIF3-f), identified in C2C12 ([440, 441], reviewed in [108]). This suggests that MAFbx may be an initiator, neutralizing a few specific muscle protecting factors, belonging to multiple metabolic pathways. Because MAFbx neutralizes a translation initiation factor, it has been suggested that MAFbx relative importance is augmented in conditions where muscle loss relies more on loss of protein synthesis. In contrast, two-yeast hybrid experiments revealed two large classes of well-expressed putative MuRF1 substrates, myofibrilar components, including titin, nebulin, troponin-I, troponin-T, myosin light chain 2, and components of ATP-generating machinery, including NADH dehydrogenase 1a, NADH-ubiquinone oxidoreductase, pyruvate dehydrogenase, and ATP synthase beta-subunit[442]. The loss of the latter, but not the former group, was confirmed in the transgenic mouse overexpressing MuRF1[443]. In vitro, MuRF1 ubiquitinates myosin heavy chains[444] and actin[445]. MuRF1 is therefore a more plausible effector of bulk protein degradation, as it is a better fit for the “undiscriminating” proteolytic machine predicated by Goldberg.

Studies on cultured myotubes confirm that GC induce the two atrogenes directly, without a third-party organ mediation[446]. The sparing of MuRF1 knockout mice revelats its central role in GAML[432]. A sizable body of indirect evidence suggests that the main positive regulator of atrogenes in GAML are FOXO transcription factors. These transcription factors are reliably upregulated by Dexa in muscle. In addition, Dexa inhibits their kinase, Akt, thus protecting them from ubiquitination and degradation. Atrogenes are also modulated by myostatin, through SMAD3 transcription factor, and by AMPK. Transgenic mice with a defective GR still exhibit upregulated atrogenes in denervation and fasting. Therefore, atrogenes integrate a multitude of GC-activated and GC-independent metabolic and hormonal signals (discussed in dedicated sections).

A series of less investigated, Foxo-independent factors lend support to Dexa’s upregulation of atrogenes. The E3 ligase TRAF6, whose expression is increased by Dexa, appears necessary for GAML and atrogene upregulation[447]. In cultured cells, Dexa upregulates the nuclear cofactor p300[448]. Dexa also upregulates acetylation and nuclear translocation of C/EBP β, in a p300 dependent-manner[429]. Because atrogenes promoters contain putative binding sites for C/EBP transcription factors, p300 was hypothesized to be yet another mechanism by which GC stimulate atrogenes. Interestingly, p300 is a histone acetyltransferase (HAT). Dexa increases HAT activity and reduces HDAC activity in muscle[449]. Because HDAC 3 and 6 are repressed by Dexa and because trichostatin A, an HDAC inhibitor, upregulates MAFbx, it has been hypothesized that Dexa acts by increased histone acetylation. The studies are interesting opening move in the field of epigenetics, and should be followed up with identification of specific chromatin loci and / or extranuclear proteins whose acetylation is modulated by Dexa. The importance of histone acetylation in in vivo GAML is still to be determined.

In conclusion, explant models suggest that GAML is, to a wide extend, the result of upregulation of the proteasome-ubiquitin system. While two important E3 ligases have been identified, the absence of specific anti-MAFbx and anti-MuRF1 antibodies prevents the study of their protein flux and their intracellular localization[108]. Judging by the significant residual atrophy after depletion or inhibition of the atrogenes, other mechanisms, within and outside the ubiquitin-proteasome system, are co-opted by Dexa in GAML. Next sections will describe the various pathways that collaborate to stimulate the ubiquitin-proteasome system. Each of them has proteasome-unrelated side effects, including activation of other proteolytic and anti-translational pathways.

## Glucocorticoid-induced loss of sensitivity on the IRS - Akt axis

Surprisingly, the most comprehensive account on GC-induced changes on protein metabolic regulation comes from the study of glucose metabolism changes. Dexa causes systemic insulin resistance, manifested as uncompensated hyperglycemia[412]. On one hand, Dexa induces concerted catabolic changes which eventually converge to hyperglycemia, and consequent hyperinsulinemia. Together, these yield a higher index of homeostatic model assessment - insulin resistance (HOMA-IR)[450]. On the other, Dexa-induced hyperinsulinemia fails to trigger its typical anabolic program in muscle. For example, Dexa overrides insulin to reduce muscle glucose uptake[451]. Dexa does not alter expression and activity of hexokinase, and of glucose transporter (GLUT) 4 expression[452], but inhibits GLUT4 recruitment to the cell membrane[451, 452]. Because translocation of GLUT4 in response to insulin is fundamentally dependent on

Akt[453], its suppression by Dexa is evidence for Dexa-imposed Akt inactivation. But in normal muscle, hyperinsulinemia causes Akt activation. Therefore, there is a disconnection between between the circulating anabolic insulin signal and the catabolic state in the sarcoplasm.

Insulin or IGF-I, in concentrations close to the physiological levels, increase glucose uptake about 10-fold[451]. The fact that there is significant leeway for amplification of anabolic pathways demonstrates that, at basal state, muscle Akt-mediated signals are diminished. Unsurprisingly, investigations of muscle in basal state have been prone to fail to observe changes in Akt-related signals. Moreover, studies on the Akt pathway are conceptually challenged by the low basal activity. In order to prove causality by downstream inhibition, one must reverse that inhibition, and establish that the final effect is lost. Chemical reversal of inhibition is rarely an option. In the case of Akt, genetic reversal of inhibition, that is, overexpression studies, have minute chances of matching wildtype basal level. Most often, overexpression of IGF-I, IGF-1R, Akt, or other mediators of

GAML overcompensates Dexa’s effect, and causes overriding, possibly non-specific and irrelevant, hypertrophy.

In contrast, Dexa co-administration blunts muscle anabolic and anti-catabolic response to exogenous supplementation with either insulin or IGF-I, at the levels of protein and glucose metabolism[454, 329]. Typical reductions in metabolism insulin sensitivity are between 40% and 80%. Most of our knowledge about Dexa effect on the Akt axis is derived from studies of muscle stimulated with pharmacological doses of insulin or IGF-I, each with their own risks of inducing experimental artifacts.

Dexa extinguishes the insulin signal very early. By itself, Dexa treatment does not change basal levels of autophosphorylated insulin receptor in absolute or stoichiometric terms[450, 455, 456]. Similarly, Dexa alone does not change the basal level of phosphorylated IRS. However, in response to insulin, mice treated for 5 days with 1 mg/(kg d) Dexa exhibit only a third of the IRS 1 and 2 phosphorylation, compared to untreated animals [457]. The suppression of insulin sensitivity at IRS level has been attributed, based on observational studies, to an improved protection of IRS by calmodulin[458], to an inhibitory phosphorylation on another residue, possibly by PKC[459, 456], or to the upregulated phosphatase C1-Ten[460]. In C2C12 myotubes, Dexa may downregulate IRS through caveolin repression[418].

The notion that Dexa interferes with IGF-I signaling suggested that GAML also attenuates downstream MAPK response (reviewed in [461]). Activated receptors for growth factors, such as IGF-1R, phosphorylate and assemble a transduction complex, including Src homology 2 domain containing (Shc) and Growth factor receptor-bound protein 2 (Grb2). Dexa reduces insulin’s ability to cause Shc phosphorylation and association with Grb2[462]. Paralleling the findings in sugar metabolism, Dexa has no detectable effect in basal state. Further downstream, the complex including Shc and Grb2 binds and activates Son of sevenless homolog (Sos), the GTPase exchange factor for Ras. Ras-GTP activates a cascade of kinases, including Raf, Mek, and Erk, eventually leading to cell proliferation. A report describes that Dexa upregulates phosphorylation of Mek and Erk in diabetic rats[427]. Others describe increased Erk phosphorylation in L6 myotubes during acute Dexa[463]. Nevertheless, no reports describe such changes in wild-type animals, which still undergo GAML. Measurements of changes in activation of another MAPK, p38, are contradictory[464, 465]. Together with the limited role of hyperplasia in adult muscle, available evidence suggests that MAPK cascades do not mediate GAML. More to the point, absence of sizable changes in the MAPK cascades reinforces the idea that insulin and IGF-I signaling is extinguished at IRS level.

In addition to interference at IRS level, Dexa hinders the recruitment of PI3K to the membrane complex containing insulin receptor and IRS1[450, 457]. Mutagenic studies demonstrate that GR’s transcriptional and DNA binding domains are required, in order for it to inhibit p70-S6K[466]. In the current model, p70-S6K is downstream of IRS and PI3K, suggesting that Dexa inhibitory action is a transcriptional effect. Two mechanisms have been found in cell culture models. One asserts that PI3K catalytic subunits are sequestered in the cytosol by an glut of its regulatory unit, p85 α[450, 467, 468], whose expression is upregulated in vivo by Dexa[435]. The other contends that activated GR binds p85 in a competitive manner, thus displacing it from IRS1[469]. This non-transcriptional effect has not been explored in GAML in vivo, but is supported by the unusual persistence of IRS1-PI3K complexes during diabetes in

GR knockout mice muscle.

The formation of a membrane complex including the transmembrane

receptor, IRS, p85 and, PI3K catalytic subunit, p110 (reviewed in [470]). The latter acts on the membrane lipids to synthesize 3-phosphoinositides. In muscle cell lines treated with Dexa in vitro, the depletion of membrane-bound PI3K and of 3-phosphoinositides leads to lower activation of 3-phosphoinositide-dependent protein kinase 1 (PDK1), and lower phosphorylation of the latter’s substrate, Thr 308 on the activation loop of Akt. While Thr 308 phosphorylation is not detectable in rat muscle in basal state, its sensitivity to insulin is depressed by Dexa[471].

In other systems, the other relevant Akt phosphorylation site, Ser 473, is paralleling the changes in Thr 308. Even in GAML, Dexa lowers insulin’s ability to upregulate Ser 473 phosphorylation[471, 457, 472, 399]. However, the effect of Dexa on Ser 473 phosphorylation in basal state is debated. Until about a decade ago, Akt phosphorylation was identified by changes in electrophoretic mobility. With that method, in basal conditions, in rat muscle, Ser 473 phosphorylation is either undetectable or below the dynamic range of immunoblot[471, 457, 472, 399]. Even in cultured cells, Akt phosphorylation is too low to be quantified[473, 446, 474]. To establish whether Dexa has an effect on Ser 473, one had to use an insulin or IGF-I challenge. Lately, the use of phospho-specific antibodies brought about a series of reports where Ser 473 phosphorylation appeared measurable[475, 476, 477]. These reports also describe that Dexa achieves a reduction in Ser 473 phosphorylation in basal state.

The relationship between Ser 473 and Thr 308 phosphorylation is a subject of ongoing research. Ser 473 phosphorylation is stimulated by growth factors, through mTOR complex 2, and through mTOR-independent,

3-phosphoinositide-stimulated mechanisms[478]. In vitro manipulations indicate that the two sites synergize for maximal Akt activation[479]. The interdependence between Ser 473 and Thr 308 is illustrated by the fact that non-mutagenic in vivo experiments rarely describe Akt activation without phosphorylation at both residues (reviewed in [480]). This correlation may be the result of an initial priming event, conditioning a non-limiting second phosphorylation.

Mutagenic abolition of Ser 473 phosphorylation abrogates Akt activity on substrates belonging to the FoxO class, but does not change Akt ability to phosphorylate tuberin (Tuberous Sclerosis Complex 2; TSC2) and glycogen synthase kinase 3 (GSK3)[481]. Such changes in substrate imply that Ser 473 controls specificity. Nevertheless, similar results would be seen if Ser 473 alters solely Akt activity, but FoxO substrates are scarcer compared to tuberin and GSK3. In either case, because Foxo is at the core of atrogene regulation in muscle atrophy, Ser 473 status has been of greater interest for GAML studies.

More evidence for Dexa-induced inhibition of Akt comes from its substrates, such as Ser 9 on GSK3-β. Chronic Dexa reduces GSK3-β phosphorylation at Ser 9, thus leading to GSK3-β activation [471]. The activation of GSK3-β in GAML is confirmed by increased phosphorylation of its substrate glycogen synthase at Ser

645, 649, 653, 657, and decreases intramuscular glycogen synthesis rate[482, 483, 471]. Among the many substrates of GSK3-β, the subunit ε of the eukaryotic initiation factor 2B (eIF2B) may mediate an anti-anabolic effect[484]. In cultured myotubes, GSK3-β knockdown reduces Dexa ability to upregulate MAFbx[485]. While inactivation of GSK3-β is typically attributed to latent Akt inhibition, the Ser 9 site is also a target for ribosomal protein S6 kinase, 90 kDa (p90-RSK), serum and glucocorticoid-inducible kinase-like kinase (SGKL), and p70-S6K[486, 487, 488]. The role of GSK3-β in muscle atrophy was briefly the subject of research at the turn of the century, in the context of a hypothesized atrophic mechanism involving calcineurin and nuclear factor of activated T-cells (NFAT). Since then, evidence that calcineurin is not involved in muscle anabolism[489] led to an unjustified neglect of GSK3-β.

In conclusion, the muscle response to GC is associated with signaling and glucose metabolism changes consistent with IRS and Akt loss of sensitivity to trophic factors such as insulin and IGF-I. Interestingly, the pathway appears largely unaffected by Dexa in basal conditions, when trophic factors are withheld. Because the model of GAML centered on Akt is derived from experiments with acute states, cell models, and intrinsic anabolic interventions, it is perfectible. The best studied downstream targets of Akt are mTOR and FOXO.

## Glucocorticoid inhibition of mTOR

One of the most effective pathways for Akt to induce its anabolic program is through mTORC1 (reviewed in [490, 491, 492]). Its modulation in GAML is clearly proven by changes in its downstream targets. However, the mechanism by which mTOR is modulated by Dexa is not elucidated yet.

Cell models established that activated Akt phosphorylates and inhibits proline-rich Akt substrate of 40-kDa (PRAS40), a negative regulator of mTORC1[493]. Moreover, the same cell models showed that activated Akt phosphorylates TSC2, thus causing its sequestration with a cytosol partner,

14-3-3, and away from its transmembrane partner, hamartin (Tuberous Sclerosis Complex 1; TSC1)[494]. The destruction of TSC1-TSC2 complexes relieves the negative regulation from Rheb, a small GTPase, that can induce activation of mTORC1, manifested in amplified phosphorylation of mTORC1 substrates such as p70-S6K. Therefore, Dexa-induced repression of Akt should inhibit mTORC1. However, the only confirmation of this mechanism comes from the observation that, in L6 myotubes, Dexa stimulates the formation of TSC1-TSC2 complexes[495].

Based on upstream changes, the effect would be subtle, detectable only during hyperinsulinemia. For always-on, feeding-independent GAML to proceed, insulin-independent pathways are also required. It has been hypothesized that Akt-independent mechanisms contribute to mTOR suppression in GAML. A well-studied negative regulator of mTOR is the energy sensor AMPK[496, 497]. However, Dexa inhibits muscle AMPK phosphorylation and activity[472, 498], probably as a consequence of intramuscular ATP upregulation[499]. The surge in intramuscular ATP is caused by Dexa-induced improvements in mitochondrial function, exemplified by upregulation of cytochrome c oxidase expression and activity[500] and of Na(+)-K(+)-ATPase expression and maximal activity[501]. Chronic Dexa or corticosterone do not alter Liver Kinase B1 (LKB1)[472, 502], the main AMPK kinase, indicating that energy sensing employs other, unknown mediator. This body of evidence shows that in GAML, mTOR suppression does not employ AMPK. Moreover, GAML does not involve mitochondrial dysfunction.

Another putative mTORC1 inactivating mechanism is centered on the stress sensor REDD1. Dexa upregulates REDD1 expression in muscle[435].

Demonstrating its key role, genetic depletion of REDD1 abolishes GAML and

Dexa-induced myotube atrophy[495, 503]. In cell culture, REDD1 interferes with 14-3-3 in order to release TSC2, restore TSC1-TSC2 complexes, and eventually inhibit mTORC1[504]. Moreover, REDD1 is in epistasis with AMPK[505], at times overriding its action on mTOR. Interestingly, a REDD1-based mechanism could explain the few observations on mTORC1 changes just as well as a

Akt-centered model would.

One way to asses mTORC1 activation is to observe its enzymatic activity. In contrast to Akt, phosphorylation of mTORC1 substrates, 4E-BP (Thr 37/46) and p70-S6K (Thr 389), is measurable in rat muscles in basal state, with acute Dexa causing a reduction of their phosphorylation[506]. Chronic Dexa changes 4E-BP and p70-S6K phosphorylation in a pattern that mirrors Akt activation, with no detectable Dexa effect on their basal phosphorylation, and a Dexa-induced loss of sensitivity to insulin[507, 454]. Conversely, insulin sensitivity at 4E-BP and p70-S6K is enhanced by removal of endogenous GC, through adrenalectomy[508, 509].

Another way to gauge the role of mTORC1 is to use its inhibitor, rapamycin. One group reported that Dexa-induced decrement in protein synthesis is unmodified, in absolute terms, upon rapamycin co-administration[454], suggesting that mTOR signaling is dispensable. In a C2C12 microarray study, interference of IGF-I transcriptional program with PI3K inhibitor was virtually identical to the interference induced by rapamycin[510], suggesting that, on the contrary, mTOR is indispensable for PI3K effects. Most likely, both hypotheses are based on reductionist models that do not reflect true in vivo phenomena.

Diverging from in vivo studies, some studies showed that L6 and C2C12 myotubes exhibit Dexa-induced downregulation of p70-S6K Thr 389 and 4B-EP phosphorylation and activities, even in the absence of an anabolic stimulus [511, 435, 473]. It is unclear whether the difference between in vivo and in vitro studies of mTORC1 substrates is caused by the use of different anabolic stimuli.

Whereas in vivo studies use insulin to elicit an Akt response whose modulation by GC is measurable, in vitro studies commonly use IGF-I (for example, [446]). One reason could be the remarkable hyperplastic abilities of insulin[512], which may perturb experiments where proliferating myoblasts were not actively eliminated.

While Akt repression is consistent with the induction of the ubiquitin-proteasome system via FOXO, its side effect, mTOR repression, is expected to lead to simultaneous downregulation of protein synthesis via 4E-BP (discussed later). Discovery of novel mTORC1 substrates such as UNC-51-like kinase 1 (ULK1)[513] led to speculations about additional contributions to the GAML phenotype from autophagy. In vivo evidence for mTOR-mediated autophagy upregulation in GAML is indirect (discussed in a later section).

Based on p70-S6K divergence, it appears that GC causes Akt-mediated mTOR modulation, in contrast with denervation atrophy, where Akt is downstream of mTOR. However, an unreplicated recent study opens the perspective for a paradigm-overturning situation. In an acute study of Dexa on female rates, Britto and colleagues confirmed most of the current model[503]. Dexa repressed mTORC1 activity, as indicated by lower phosphorylation of its substrate 4E-BP1. The repression of mTORC1 was correlated with an reduced inactivation of the negative regulator PRAS40. In contrast, and unexpectedly, acute Dexa increased TSC2 phosphorylation, indicating that Dexa induces substrate selectivity changes in Akt. As discussed earlier, such Akt selectivity alterations are attributed to dissociation between Ser 473 and Thr 308 phosphorylation state. Indeed, Britto and colleagues found that, after acute Dexa, phosphorylation at Ser 473 increases, while at Thr 308 it decreases. This is a rare, although not unique, account of Akt hyperphosphorylation in GAML. In each case, these findings may be explained by the particular conditions of the unique experiment. Here, proteolysis was not unaltered, indicating that acute Dexa is not a true model of GAML.

Britto and colleagues refined their model by observing the effect of Dexa in REDD1 knockout mice. REDD1 depletion abolishes GAML, although atrogenes are still upregulated. On the other hand, REDD1 depletion prevents Dexa’s hypophosphorylating action on 4E-BP1 and p70-S6K. These findings place mTORC1 inhibition at the core of GAML. Britto and colleagues propose a model of GAML where Ser 473 hyperphosphorylation is caused by an mTORC1 feedback loop being severed by Dexa. Upstream of mTORC1, REDD1 knockout mice lost hypophosphorylation on PRAS40 and of Akt on Thr 308. While Akt remains upstream of mTOR and vital for GAML, Britto’s model discards phosphorylation of Akt at Ser 473 and of TSC2 from the rapid REDD1-dependent response to Dexa. Multiple studies in transformed cells confirm the role of Thr 308 in REDD1 repressive action on mTOR, while proposing different intermediaries ([514]; reviewed in [515]).

The most robust inducer of REDD1 is hypoxia, through HIF transcription factors[515]. As described in the dedicated section, HIF1a also induces VEGF, which in turn has a myoprotective action. But in GAML, VEGF is downregulated[516], suggesting that HIF1a is not involved in REDD1 upregulation. The mechanism by which Dexa stimulates REDD1 remains to be studied. Candidates include ATF4 and NFAT, discussed in other sections of this chapter.

While it is certain that GAML is correlated with mTORC1 downregulation, it is still unclear how mTORC1 repression is achieved, and what is its relative importance. Based on the evidence published until now, the recent change of paradigm which placed p70-S6K at the forefront of denervation atrophy appears unlikely in GAML. For now, our understanding of GAML places Akt firmly upstream of mTORC1. Better transgenic models, a deeper understanding of mTOR pathways and more in vitro / recombinant experiments are required for a

definitive conclusion.

## Glucocorticoid activation of Foxo transcriptional program

The evidence for the role of Foxo in GAML is not as direct and compelling as that gathered in denervation atrophy. Given the partial redundancy between the three regulatable Foxo transcription factors, only a triple knockout will give the true measure of their relative importance. Nevertheless, an impressive body of indirect evidence supports their involvement in GAML.

The importance of Foxo transcription factors is supported intuitively by their role as integrators of multiple atrophy signaling pathways. Multiple mechanisms converge to induce Foxo upregulation in GAML.

First, chronic Dexa doubles Foxo expression[435]. Foxo1 promoter contains a GR response element[517]. In C2C12 myotubes, the response is biphasic, combining a short-lived ample and rapid increase, with a gentler long term augmentation[436]. Foxo promoters contain GR-binding sites[518]. In vitro,

Foxo induction by GR is facilitated by the histone acetyl transferases p300 and CREB-binding protein, which are independently upregulated by Dexa[449]. As a consequence, GR knockout reduces expression of FOXO1 and FOXO3a in muscle[433].

Second, Foxo transcription factors are thought to be potentiated by Dexa through post-translational means involving the earlier described Akt pathway (reviewed in [519]). Akt phosphorylates Foxo1 at Tyr 32 and Ser 253, thus creating binding sites for 14-3-3. Phosphorylated Foxo transcription factors are exported from the nucleus, and eventually marked for ubiquitin-proteasome-mediated degradation. It is believed that the Dexa-induced impairment of Akt contributes to an increase in their activity in GAML. Given that phosphorylated Foxo is degraded and is elicited by a low-activity Akt pathway, Foxo phosphorylation changes may be undetectable in basal state. In C2C12 myotubes, Foxo phosphorylation downregulation in response to Dexa is more readily detected[383, 520]. A few groups reported reductions in Foxo phosphorylation by chronic Dexa at basal state in vivo[521, 475, 522]. Two out of the three referenced studies also report an unusual activation of Akt in basal state, indicating that they may be reflective of special conditions (40 days treatment, the use of male levator ani, and so on). As mentioned in previous sections, an ongoing debate surrounds the effect of Dexa on Akt Ser 473, which alters its ability to phosphorylate FOXO.

As with mTORC1, the Akt pathway may be more relevant in fed state.

Third, PGC nuclear cofactors may facilitate the Dexa-induced Foxo surge. Loss of sensitivity to insulin causes PGC-1 repression in muscle[523]. In particular, Dexa represses muscle PGC-1α[464, 522] and PGC-1β[524]. In cultured myotubes, PGC-1β overexpression and knockdown cause changes in the reverse direction for Foxo3a and atrogenes’ mRNA. Because PGC-1α overexpression drives conversion of fast to slow twitch fibers[352], it has been speculated that differences in PGC-1α levels make different types of muscle fiber more or less sensitive to Dexa[384]. A whole higher level of regulation of is yet to be studied, with PGC-1α splicing dictating its role more. For example, isoform 1, and not 4, of PGC-1α is correlated with muscle hypertrophy[525].

Fourth, indirect evidence links Foxo1 to PPARβ/δ. Dexa stimulates simultaneously PPARβ/δ DNA binding and Foxo1 acetylation[526]. PPARβ/δ inhibition reduces GAML and, at the same time, prevents Dexa-stimulated acetylation of Foxo1. While the direct effects of this post-transcriptional modification is not elucidated (reviewed in [527, 528]), it was speculated that PPARβ/δ changes induce physiologically relevant alterations in Foxo1.

Finally, miR-182, whose expression is inhibited by Dexa, is a negative regulator of Foxo3[529]. Other mechanisms, such as AMPK, have been hypothesized, based on the latter’s ability to upregulate atrogenes.

The net result of these mechanisms is that Dexa treatments upregulates nuclear Foxo[464]. More convincing evidence towards Foxo activation comes from the concordance in its downstream effects. For example, Dexa increases muscle glutamine synthetase activity[530]. Glutamine synthetase promoter contains a FOXO response element. Dexa-induced expression of muscle glutamine synthetase is absent in Foxo1 knockout[531].

Manipulations of Foxo have supported their role in GAML. Transfection with dominant negative Foxo3A partially prevents diameter reduction in disuse of rat soleus and Dexa-treated C2C12 myotubes[520, 532].

Among the Foxo-induced genes, a special importance was given to the E3 ligases associated with muscle loss, MAFbx and MuRF1[446]. Both atrogenes contain FOXO-binding regions in their promoter[520, 533, 433]. These Foxo-responsive elements are in close proximity of SMAD3-binding elements, in a shared response element which facilitates synergistic interactions between the two classes of transcription factors[534]. MuRF1 promoter facilitates another synergistic interaction, between FOXO1 and GR[433]. A positive feedback loop links MuRF1 and FOXO1, as indicated by downregulation of p85 and FOXO1 in late GAML in the MuRF1 knockout[417].

In both atrogenes’ promoter, a KLF-15 response element is located, near the FOXO binding site[535]. KLF-15 is a direct transcriptional target of GR, and interferes with mTOR signaling. A complex network is centered on KLF-15, which upregulates expression of FOXO transcription factors, and synergizes with them to upregulate atrogenes. Overall, atrogenes appear to be under a strong control of Foxo, favored by concomitant activation of synergistic factors. Indeed, dominant negative Foxo3A transfection in rat soleus leads to loss of atrogene upregulation[536]. In cultured myotubes, where transfection of constitutively active FOXO3a upregulates MAFbx[520], while knocking down FOXO1 causes a reduction in the atrogene response to GC[537].

A majority of the effectors described in this chapter appear to be under the control of Foxo pathway. In addition to atrogenes, Foxo targets include 4E-BP1, cathepsin L, and the effector of denervation atrophy, Growth Arrest and DNA Damage 45 (Gadd45)[538].

## Glucocorticoid activation of myostatin

In rodents, Dexa upregulates myostatin, a strong negative regulator of muscle mass, in the atrophying muscle[539]. In myostatin knockout mice, Dexa induces most of its transcriptional program, including modulations of IGF-I, MuRF1, MAFbx, FOXO3A, but Dexa-induced muscle loss does not occur [423]. These findings suggest that myostatin is an important mediator of Dexa.

Myostatin’s promoter contains GR response element[539], a FOXO responsive element[540], and a CCAAT sequence[541], all known to be stimulated by Dexa. Myostatin expression is also upregulated by the histone methyltransferase SMYD 3, which in turn is upregulated in muscle by Dexa[542]. The stability of the myostatin transcript is improved by Dexa-exerted repression of its negative regulators, miR-27[543].

Glutamine supplementation reduces GAML and myostatin expression in muscle[544]. This finding suggests that, in the context of GAML, myostatin may double as a nutrient sensor.

Myostatin’s effects are manifold, and not completely understood. For example, while myostatin inhibition in utero leads to doubling of muscle mass in adult animals[225], its overexpression in adult animals causes a more moderate effect[545], indicating that its effect is based on hyperplasia- and hypertrophy-based pathways of muscle growth. In explanted adult mouse myofibers, where proliferation has been minimized, expression of a dominant negative form of activin A receptor, type IIB (ActRIIB), the receptor for myostatin, causes an increase in CSA, which is halved by rapamycin[546]. Therefore, proliferation-independent mechanisms of myostatin-induced atrophy may include both mTOR-dependent and -independent pathways. The extensive, multifactorial action of myostatin on muscle suggests that it may override any atrophic stimulus, in non-specific ways. Recent molecular studies endeavored to establish whether GAML-associated myostatin modulation is physiologically relevant, or only a side effect.

A series of experiments describe Akt inhibition in response to myostatin treatments. In vivo electroporation with myostatin-coding plasmids leads to 10% loss in tibialis anterior mass in the transfected muscle, in the absence of any changes at the level of atrogenes[547]. The investigators behind this study performed an analysis of the mTOR pathway, which found loss of phosphorylation on tuberin and p70-S6K, and concluded that myostatin affects protein metabolism. A simpler explanation would have taken into account the regeneration in the scarred electroporated muscle, which should have contributed a high number of proliferating cells. Another provoking study investigated human myotubes differentiated from cultured myoblasts. In this model, myostatin treatments lead to reductions in diameter and atrogene downregulation [548]. The authors of the latter report explain their observations by demonstrating that myostatin reduces myoblast fusion and differentiation. While interesting in context of muscle development and regeneration, such experiments cannot replicate adult GAML. Moreover, both reports describe even in the myostatin-free controls, an unusual Akt phosphorylation. Most of their observations would be consistent with the well-documented myostatin-induced impairment of muscle regeneration[549, 550].

The study on human myotubes[548] showed that exogenous myostatin downregulates muscle atrogenes. In contrast, GAML is characterized by an upregulation of atrogenes. Moreover, the upregulation of atrogenes in GAML is attributed, in part, to Akt inhibition. Therefore, a myostatin - Akt axis does not seem to underlie GAML.

Myostatin is particularly adept at modulating the fusion of myoblast to myotubes[551]. Myostatin’s action appears more relevant for processes where myoblasts or satellite cells are involved, such as muscle development and regeneration (reviewed in [552, 553]). But normal adult muscle maintenance involve to a small degree satellite cells. Loss of their contribution to muscle maintenance, which is supposedly a major effect of myostatin, may contribute to, but not command adult GAML. In contrast, there is no evidence that myostatin upregulates catabolic processes within GAML. Even in other models of atrophy, such as microgravity, myostatin upregulation occurs while 3MH excretion is unchanged[554]. Myostatin’s relative importance for GAML remains unclear. The typical experiments, using qualitative alterations of myostatin, may override any endogenous mechanism, and yield non-specific anti-atrophic effects.

## Glucocorticoid modulation of protein synthesis

Goldberg’s[372] 1969 study on cortisone-induced atrophy of rat muscle brought one more important theme for the field of GAML. In that experiment, the specific activity, that is, the ratio between tracer and total protein, was the same in GC-treated muscle as in control animals, despite decreased muscle mass and decreased total tracer in the GC-treated group. Goldberg conjectured that an equal rate of protein synthesis in the two groups would have caused a faster dilution of the tracer in the GC-treated muscle. Failing to find that was interpreted as evidence for decreased protein synthesis in cortisone-treated muscle. This was an elegant way of bypassing the denominator effect from the typical experiments, where synthesis rate would be measured over hours, in muscles that already underwent atrophy in the prior days. In these cases, a loss in protein synthesis rate might be underestimated or even factored out, after normalization to a lower muscle mass.

As in human studies, the role of protein synthesis modulation during chronic hypercortisolism is still debated, because chronic Dexa effect is superimposed by acute effects. A single 20 mg/(kg d) triamcinolone dose nearly halves incorporation of labeled precursors in rat muscle protein at 8 hours, yet has no effect after 24 hours[555]. Similar acute effects may be expected from feeding, which causes fluctuations of insulin and possibly of the Akt / mTOR / 4E-BP axis in muscle.

Variability in the time from feeding and treatment to biopsy is a significant source of experimental noise, which must contribute to inconsistencies in literature. For example, some of the most exhaustive studies of protein metabolism in GAML came from Grizard laboratory. In two studies, they found that protein synthesis rate was unchanged after 0*.*5 mg/(kg d) Dexa for 6 days, compared to pair-fed animals, in the epitrochlearis muscle[411, 556]. In two other studies, assaying other glycolytic muscles or quadrupling the Dexa dose, led to observable reductions in protein FSR[557, 454]. In cell cultures, Dexa inhibition of protein synthesis should be more readily detectable, given that, in vitro, Dexa affects 4E-BP phosphorylation directly, even in basal state[473]. However, studies are similarly equivocal. In the most glaring example, the same group, using the same methods and working on the same L6 line, found that protein synthesis is

“not altered” by Dexa in one experiment, and decreased in another[406, 558].

Even when detected, the amplitude of changes in protein synthesis in vitro[558, 559, 560] and in vivo[329] is lower than that on protein degradation, when Dexa is given at the dose that causes muscle loss. In contrast, doses of Dexa below the apparent EC50 are unable to change protein degradation rate, while still lowering protein synthesis rate[336].

In the MuRF1 knockout mouse, Dexa fails to change FSR[432], suggesting a temporal order, with proteolytic processes preceding and initiating changes in protein translation.

Suggestive data for protein synthesis modulation in GAML come from various molecular correlates. For example, acute GC administration causes a reduction in the proportion of polyribosomes[561]. The proportion of polysomes recovers to basal level in less than 24 hours, when the GC is short-acting prednisone[407]. In contrast, polysome downregulation lasts about two days after Dexa. No published accounts describe polysome profile changes after longer GC exposure.

Dexa has a dual effect on 4E-BP, the inhibitor of protein synthesis that binds and inhibits the eukaryotic initiation eIF4E. First, Dexa upregulates 4E-BP mRNA[435]. It was long believed 4E-BP induction is direct, and based on a Foxo response element in 4E-BP promoter[562]. As of 2015, this response element is proven only in Drosophila. The only replication in mouse cultured cells was withdrawn during the preparation of this thesis, for unspecified reason. Second, Dexa may regulate 4E-BP through post-translational phosphorylation. As shown earlier, Dexa reduces mTOR-effected 4E-BP phosphorylation sensitivity to insulin, which is expected to cause reductions in protein synthesis rate mainly after feeding. The study of mTOR action is challenged by the existence of multiple phosphorylation sites, inability of antibodies to discern them, and inability of rapamycin to inhibit mTORC1 action on 4E-BP1[563].

Dexa upregulates mitogen-activated protein kinase–interacting kinase 2 (MNK2) expression[564]. Dexa reduces eIF4G Ser 1108 phosphorylation in wild type mice, but not in MNK2 knockout mice. This posttranslational modification correlates with nutrient availability, but its role in GAML or in general is not known (reviewed in [565]).

In C2C12 myotubes, GAML may repress protein synthesis by MAFbx-initiated neutralization of eIF3f[441]. Overexpression of eIF3f causes hypertrophy, and eIF3f knockout induces atrophy, supporting the hypothesis that loss of an initiation factor causes reductions in the rate of protein synthesis. Moreover, further data from the same group shows that loss of eIF3f impairs the ability of mTOR to bind and phosphorylate p70-S6K, suggesting that eIF3f pro-anabolic action is more complex[566]. The role of eIF3f in GAML has not been confirmed in vivo.

Canonical control of protein synthesis includes translational derepression, a cytosol-based mechanism for sensing amino acid starvation (reviewed in [567]).

Relative lack of amino acids enriches uncharged tRNA, which bind and activates General Control Nonderepressible 2 (GCN2)[568]. Activated GCN2 phosphorylates eukaryotic initiation factor 2 (eIF2) at Ser 51 on its α subunit, leading to the formation of an inactivating complex with eIF2B[569]. The inactivation of eIF2B, the guanine exchange factor for eIF2, leads to general translation shutdown[570]. The lack of eIF2-GTP complexes leads to start codon skipping, which, for transcripts including multiple open reading frames (ORF), determines extraordinary translation from downstream start ORFs. In eukaryotes, a physiologically relevant downstream ORF is Activating transcription factor 4 (ATF4), which upregulates genes involved in transport of essential amino acids and synthesis of non-essential amino acids, such as asparagine synthetase[571]. Overall, the translational derepression pathway is a mechanism for inhibiting protein synthesis, initiated by apparent depletion of free amino acids, and leading to ATF4 upregulation.

The study of translational derepression in mammalians is still in its beginnings. The ATF4 knockout mouse has normal weight, but exhibits some sparing from muscle atrophy in response to starvation[572]. ATF4 knockout mice exhibit a normal atrogene response to starvation, indicating that ATF4 is part of a novel atrophy pathway. The only study that measured chronic Dexa effect on phosphorylation of eIF2 reported negative results[123]. However, Dexa abrogated amino acid infusion ability to reduce eIF2 phosphorylation, a situation reminiscent of Dexa’s action on Akt. In a fibroblastic cell line, Dexa upregulated ATF4 translation as long as insulin was withheld from the medium[573].

Unexpectedly, in C2C12 myotubes, ATF4 protein levels are upregulated by insulin in a rapamycin-dependent manner[573]. A plausible explanation is that ATF4 is upregulated by apparent amino acid deficits, including cases when mTOR-stimulated protein translation depletes the free amino acid pool.

Changes in 4E-BP1 and perhaps some other pathways regulating protein synthesis are induced by Dexa in a manner consistent with GAML. The mild alterations of translation molecular markers correlate with the moderate loss in protein synthesis rate. While the overall effect is not negligible, its reduced amplitude and its downstream relation to MuRF1 indicate that anabolism adjustments play a complementary role to the activation of the ubiquitin-proteasome system in GAML.

## The effects of glucocorticoids on autophagy

Proteasome inhibition does not abolish completely Dexa-activated proteolysis, indicating that some other catabolic mechanisms must be involved. Based on newly discovered role of mTOR as a major modulator of autophagy, the second most important effector of GAML was presumed to be autophagosome. Moreover, in L6 myotubes, 100 µM chloroquine or 200 mM E-64 reduce the rate at which Dexa amplifies proteolytic tracer release[424].

A series of molecular markers further support the idea that autophagy is upregulated by Dexa. In C2C12 myotubes, Dexa may induce the formation of double-membrane autophagic vesicles, although evidence is limited to unquantified micrographs[574].

Dexa reliably upregulates the family of lysosome proteases known as cathepsins. In vivo, Dexa doubles the lysosome proteases cathepsin L and D

[411, 575, 436]. In L6 myotubes, Dexa upregulates cathepsin B[424].

A new modality for investigating autophagy hinges on one of the longest-living markers on its surface, LC3. Transgenic mice with expressing LC3-GFP have been developed, and are regularly used for tracking autophagosomes. There are no reports of this model being used in the study of GAML. Alternatively, endogenous LC3 is tracked through immunofluorescence microscopy. Some reports describe an accumulation of punctate LC3-containing structures in L6 myotubes treated with Dexa for 6 hours[463]. Another way of tracking LC3 is based on immunoblot. A lipidated form of LC3, termed LC3-II, migrates faster than its precursor, LC3-I, during electrophoresis. Various indices, such as the amount of LC3-II, or the ratio between LC3 electrophoretic forms, are used for estimating the number of autophagosomes (reviewed in [576]). LC3-II is enriched in C2C12 myotubes overexpressing LC3[574] and in L6 myotubes[463]. Similar to the microscopy experiments, these reports use acute Dexa treatments. Moreover, a time course reveals that in L6 myotubes, LC3-II peaks at 6 hours and is extinguished at 24 hours after Dexa administration[577]. It appears that in vitro Dexa causes a rapid accumulation of autophagosomes, which are dismantled at a lower pace, as they fuse with lysosomes, and their content is cleared by proteolysis.

In vivo, acute Dexa causes accumulation of LC3-II protein[503]. Recently, the enrichment of LCII-B after chronic Dexa has been reported[578]. The latter report is uncommon, and may be reflecting a short interval between last Dexa dose and sample collection.

In contrast with the limited body of evidence we have for autophagy upregulation, there is a significant amount of literature describing what would drive autophagy up in GAML. In C2C12 myotubes, overexpression of a constitutively active FoxO3 or chemical inhibition of Akt leads to increased lysosomal proteolysis[383]. As mentioned earlier, mTORC1 is a negative regulator of autophagy, whose inhibition in GAML is hypothesized to stimulate autophagy. Indeed, acute Dexa administration reduces ULK1 phosphorylation at Ser 575, the mTOR-specific, inhibitory site[503].

Other lines of evidence provide indirect support for autophagy upregulation in GAML. Acute Dexa downregulates p62, one of the shortest-lived markers on the autophagosome, thus suggesting that autophagic flux is upregulated[503].

Acute Dexa increases expression of the mitophagy effector Bnip3[503, 463]. In C2C12, acute Dexa upregulates expression of lysosomal markers such as autophagy-related 12 (Atg12)[529].

Given the spatial segregation between lysosomes internal and outer space, autophagy must rely on a selective mechanism for any protein that it is processing. Therefore, it is likely that autophagy acts only on a few specific, perhaps limiting, proteins. For example, Dexa-induced depletion of sialidase Neu2 is prevented by 3-MA[574].

Autophagy may play a significant role in triggering and regulating GAML.

Current evidence, based on cell culture experiments, is far from satisfactory. Given its limited amplitude in vivo, it is improbable that autophagy is responsible with bulk protein elimination.

## Glucocorticoids and other proteolytic systems

In 1986, it was discovered that proteolysis in muscle is increased when explants are soaked in 2.5 mM calcium[579, 580]. The calcium-stimulated proteolysis subsides upon co-administration of leupeptin, a wide-spectrum protease inhibitor. The discovery of a class of calcium-dependent proteases, called calpains, suggested that they might be contributing to muscle atrophy. The calpain system includes μ-calpain, which is activated by micromolar concentrations of calcium, m-calpain, which is activated by millimolar concentrations of calcium, and their inhibitor, calpastatin (reviewed in [581]). Transgenic mice overexpressing calpastatin have 30% lower loss of muscle upon unloading[582], proving that the calpain system is important in some atrophy models.

There is limited evidence for calpains involvement in GAML, beyond the experiments from 1980’s. In vivo, expression of the calcium-dependent protease m-calpain is trebled by Dexa[411]. In L6 myotubes, calpastatin overexpresion halves Dexa-induced proteolysis[583]. Still in L6 myotubes, Dexa promotes store-operated calcium entry, the mechanism by which intracellular Ca concentration is increased when ER stores are depleted[584].

It has been speculated that activations of calpains is an initial step in GAML, allowing myofibril protein to interact with MuRF1[585]. Calpains are important for some atrophy models, but their role in GAML is understudied.

Many unbiased studies found that a family of proteased, metallothioneins, are upregulated in GAML[586]. However, they contribution to GAML has not been analyzed.

With the advent of new genome technology, enzymatic and functional studies that brought the proteasome in the center of GAML have been abandoned. The example of calpains illustrates how non-transcriptional events can contribute to GAML, and testifies to a blind spot in GAML research.

## Alleviation of glucocorticoid myopathy by IGF-I

Upon finding that, in L6 myoblasts, Dexa-stimulated proteolysis is abated by co-administration of insulin, Ballard hypothesized in 1983 that Dexa acts indirectly, by depleting body’s supply of IGF-I[587]. While the use of non-fusing myoblasts is certain to introduce confounding changes in proliferation, and the equivalence between IGF-I and insulin lacks subtlety, his hypothesis captured the attention of many investigators, despite going against another theme of interest in those days, the hyperplastic synergy between IGF-I and GC[588, 589]. As described in the sections on Akt and mTOR, most of the knowledge we have about it is derived from studies of anabolism, where muscle, rather than being analyzed in its basal state, is first stimulated with insulin or IGF-I. In a sense, we know more about the interaction between Dexa and IGF-I than we know about Dexa alone. Lately, the interpretation of such experiments shifted from mechanism-explaining to a therapeutic paradigm. In mice, electroporation of IGF-I plasmid in tibialis protected solely the transformed fibers from Dexa-induced atrophy[590]. In Dexa-treated rats, co-administration of IGF-I reduces loss of muscle mass, fiber atrophy, and 3MH release[591], thus providing a blueprint for the ideal anti-GAML therapy.

Co-administration of IGF-I reverses upregulation of ubiquitin, and of proteasome subunits C2 C3, C8[592, 593], thus blunting one of the major effectors of GAML, the ubiquitin-proteasome system. But Dexa antagonizes IGF-I on many other downstream effects, such as glutamine synthesis[594].

In vivo, Dexa reduces muscle expression of IGF-I[595, 200], and possibly interferes by altering IGFBP secretion[435]. Therefore, GAML manifests as an absolute loss of IGF-I exacerbated by a downstream inhibition. Intuitively, IGF-I supplementation may provide GAML alleviation.

Despite the positive results seen in rodents (described above), very few studies analyzed molecular mechanisms by which IGF-I works in vivo. Because co-administration of GH cannot reverse muscle loss from Dexa or triamcinolone[596, 593], it is likely that, similar to Dexa, IGF-I action on muscle is cell-autonomous. Therefore, a series of studies dissected the molecular mechanisms by which IGF-I could alleviate cultured myotube atrophy. There, IGF-I alone improves protein synthesis, but has no effect on protein degradation[597]. Nevertheless, myotubes treated with Dexa, co-administration of IGF-I has an anti-proteolytic effect[597]. The reduction in proteolysis covers all domains, but, in acute settings, appears more effective in repressing lysosomal than proteasomal activity[598]. IGF-I co-administration reverses MAFbx and MuRF1 upregulation[489, 433], in a Foxo dependent manner. During co-administration, Dexa is still able to upregulate protein levels of Foxo transcription factors, but IGF-I appears to counter them by increasing their phosphorylation[446] and reducing its ability to bind MuRF1 promoter[433]. The IGF-I induced FOxo phosphorylation is consonant with upstream modulations of the Akt axis. First, IGF-I co-administration in cells treated with Dexa causes myoprotective increases in phosphorylation of Akt, GSK-3β, p70-S6K and 4E-BP1[599, 600]. Second, studies with chemical inhibitors revealed that IGF-I protective effect is mediated by Akt and PI3K[599].

From the similarity between muscle protection conferred by IGF-I and by GSK-3β inhibitors, it was speculated that IGF-I acts by inhibiting

GSK-3β[601, 476].

Muscle accretion is not completely conditioned by availability of IGF-I. Myostatin knockout mice have double muscle size, yet lower circulating IGF-I levels, compared to wild type[602]. Muscle growth and recovery is halved by rapamycin, indicating that other anabolic mediators are as important as the Akt / mTOR pathway [603, 382, 604].

Based on the current evidence, it appears that IGF-I alleviating action is not completely overlapping with the wide spectrum of atrophic actions of Dexa. Indeed, the putative mediator of muscle loss REDD1 is upregulated by acute insulin or IGF-I[511]. It is unclear whether such REDD1 upregulation in an anabolic context may refute the role of REDD1 in muscle loss, or whether, on the contrary, it explains the incomplete reversal of GAML by IGF-I.

As our understanding of GAML improved, studies on its alleviation by IGF-I are lagging. A large number of publications focus on the balance of GC and IGF-I on Akt in cultured cells. Given the reduced number of co-administration in vivo studies, our understanding of how IGF-I could alleviate glucocorticoid myopathy is incomplete.

## Alleviation of glucocorticoid myopathy by anabolic steroids

Human studies demonstrated that AAS addition to chronic Dexa benefits male adult patients, by reducing their loss of muscle and improving their quality of lifes[158]. The idea of alleviating the CS by AAS therapy came two years after the discovery of an affordable source of Testo. In the case of exogenous hypercortisolism, the same idea surfaced less than a year after the discovery of GC. Two years later, Courier and Marois report the first replication of the myoprotective effect in rats[605]. For a long time, during the second dark age of steroids, the combination of AAS and GC, with androgenic, but without anabolic potency, was investigated in the hope that it would show the way towards splitting the anabolic from the androgenic principle in Testo.

A related theoretic question was the nature of the nuclear receptor. At the time when it was not clear how many species of nuclear receptors there are, precise measurements tried to find putative interference between the various steroids, with Dexa and Testo among the most studied compounds. In vitro studies have quantified androgens ability to interfere with the binding of Dexa to GR. The Kd for GR-Dexa association is below nanomolar range[606]. In vitro Dexa doses used in muscle atrophy experiments range in the tens of hundred nanomolar. Similar concentrations are likely in the blood of mice injected with the 0.5-1 g/kg doses described before. At a concentration of 2 pM in rat skeletal muscle[607], virtually all GR should be bound to Dexa. On the other hand, Ki for Testo competing with Dexa for binding to muscle protein extracts is 10 µM[608]. While the referenced report does not distinguish non-specific binding for T, Ki is tens or hundreds of times higher than typical Testo concentrations used in literature for biological reversal of GAML. Testo binding affinity to Dexa binding sites in rat muscle cytosol is less than 100 times lower than Dexa’s affinity[609]. Therefore, barring allosteric effects, the direct competition between the two steroids remains only of theoretical importance.

The experiments in the 1980’s and 1990’s tested the interaction of AAS and

GC on the diaphragm. While these studies could not have measured the yet-undiscovered mediators of GAML, they established that myoprotection provided by AAS manifests in both muscle mass and force[610, 611, 612].

The mechanism by which AAS accomplishes muscle protection remains unclear to date. One early study found that, in vivo, Testo re-establishes the percentage of ribosomes that are involved in translation[613]. Two later studies on L6 myotubes stated that AAS cannot reverse the downregulation of protein synthesis induced by Dexa[587, 614]. A study on C2C12 myotubes also rejected an action of Testo on protein metabolism, despite a trend for improved protein synthesis and reduced protein degradation when 1 µM Testo is added to 100 nM Dexa[559]. Given the limited number of attempts, the failure of in vitro systems to replicate in vivo benefits may be ascribed to reduced sensitivity rather than to fundamental shortcomings of the in vitro model.

A few studies investigated the interaction of AAS and GC, by measuring the changes in one signal when the other is altered. An interaction at the level of receptors cannot be excluded. In skeletal muscle, GR mRNA and binding activity are increased upon castration[615, 616], suggesting a way by which castration causes muscle atrophy. Conversely, Dexa reduces the expression of AR in skeletal muscle[200]. Hypercortisolism reduces endogenous Testo levels in male rats, thus leading to ampler loss of muscle than in females, which experience Testo upregulation[413]. Sexual dimorphism in animal models confirm that males stand to benefit more from AAS therapy in GAML.

In the absence of direct interference, studies sought downstream effectors at which Testo could prevent Dexa’s program. Myostatin promoter contains putative androgen responsive elements[539]. In intact animals and even in atrophic muscle after spinal cord injury, Testo does not alter myostatin[617, 618].

Myostatin changes in co-administration of AAS and GC has not been investigated. Given myostatin’s limited role in GAML, it is unlikely myostatin repression could contribute to the ample alleviation provided by Testo.

The other effector of GAML is the Akt axis. Further suggesting a role for Akt, multiple models of hypertrophy in humans and rodents correlate with increased muscle IGF-I expression[619, 620]. However, Testo is dispensable for IGF-I upregulation during exercise-induced muscle hypertrophy[621]. Conversely, IGF-I signaling deprivation only halves T-induced muscle restoration after castration-induced atrophy[622]. In the era of immunoblot, only three laboratories published studies of co-administration of AAS and GC. All three converged towards describing an reversal of Akt pathway inhibition.

Sheng laboratory describes the effect of 10-day 1 mg/(kg d) Dexa and/or 13-day 5 mg/(kg d) T[623] on rat gastrocnemius. With these doses, Testo reliably reverses losses in body weight, muscle mass, and fiber CSA, induced by Dexa. At these doses, Testo co-administration exhibits a slight trend towards inferiority to basal levels. Molecular pathways largely confirm an Akt centered disruption of GAML. Muscle IGF-I expression is repressed by Dexa, and returned to basal level by Testo co-administration. Akt phosphorylation at Ser 473 follows a similar pattern, with Testo addition restoring what Dexa downregulates. Moreover, p70-S6K phosphorylation at Thr 389 and atrogenes correlate with Akt phosphorylation changes, indicating that Testo acts upstream of mTOR and Foxo. The only divergence from the above pattern is at GSK-3β level, where Dexa has no

effect.

Dalton laboratory account describes the effect of 8-day 600 µg/(kg d) Dexa and/or 25 mg/(kg d) T[521] on rat muscles. This study is focused on the anabolic-androgenic split, and therefore compares levator to extraperitoneal muscles. At these doses, Testo co-administration restores levator ani, but not other muscles. In levator ani, Testo co-administration upregulates IGF-I beyond basal levels, while in extraperitoneal muscles, IGF-I is only partially restored. Possibly as a consequence, Testo addition reduces atrogene expression to a larger extend in levator ani. In levator ani and C2C12 myotubes, Testo reverses losses of Akt (Ser 473), GSK-3β (Ser 9), FOXO3a (Ser 253), and p70-S6K (Thr389) phosphorylation.

The series of studies from Cardozo laboratory is the most informative. This was also the first group to show that 28 mg/(kg d) Testo reverses losses in rat gastrocnemius induced by simultaneous 700 µg/(kg d) Dexa[624]. They ascribed this alleviation to a reduction in proteolysis and atrogenes expression. In preliminary studies on L6 and C2C12 myotubes overexpressing AR, they found that Testo co-administration reverses transcriptional upregulation of REDD1 and MAFbx[625, 435].

Arguably, the most important contribution they made is the first microarray study on alleviation of GAML with T[435], using gastrocnemia treated as above. This is also the first published microarray study of GAML, generating many hypotheses still untested. Many of the genes found in the unbiased approach are consistent with the above description of GAML, including transcriptional reversal at the levels of IGFBP, IRS1, FOXO1, p85, and 4E-BP1. These changes point to a pleiotropic program through which AAS re-establish signaling on the Akt axis. The overlap is not perfect, with important presumptive effectors, including IGF-I and MAFbx, failing to pass amplitude and statistical significance thresholds.

The microarray study also describes a series of IGF-I-independent effectors. The reversal of GAML is also correlated with a repression of REDD1 to basal levels, suggesting that mTOR is also restored by Testo. Changes in B-cell leukemia/lymphoma 3 (Bcl3) and IκB indicate that GAML relies on NF-κB, and Testo is deactivating that pathway too. Testo also reversed the surge of MuRF1, proteasomal subunit D8, cathepsin L, LC3, M-type calpain, C/EBP β and δ, and metallothionein 1, exhibiting a wide-spectrum anti-atrophic action.

Dexa upregulates CD36, the fatty acid translocase[626], while Testo returns it to basal level, suggesting that GAML and its reversal have opposite effects on muscle’s ability to use lipids as fuel.

Testo reverses the upregulation of the stress sensor Gadd45, isoform β. Recently, it has been shown that Gadd45β is an autophagy blocker[627]. This suggests that in 7-day treated muscle autophagy must be downregulated, and goes against many other molecular markers that indicate the opposite.

In conclusion, the mechanisms by which Testo prevents GAML appear manifold. However, many of the putative myoprotective actions of Testo are only suggested by Cardozo’s microarray, but have never been validated.

## 3. HYPOTHESES

The example of Gadd45, at the end of the previous section, illustrates the issues GAML research is confronted with. On one hand we have macroscopic certainties, such the loss of muscle mass following GC therapy. On the other, we have invariable molecular correlates of Dexa treatment, such as augmented proteolysis, increased Foxo transcriptional activity, reduced insulin sensitivity at

Akt and mTORC1 level, and upregulation of MuRF1 and REDD1. For many of these, inhibition or interference experiments showed a reduction in muscle atrophy, but none was found to be indispensable for GAML. The hierarchy of molecular events in GAML is unclear, and lags behind progresses made in the study of denervation atrophy. Our understanding of androgen alleviation of GAML is even more incomplete.

One reason why key mediators of GC and AAS actions have not been identified is the lack of transgenic models. Muscle-localized conditional knockout experiments are needed. For this purpose, studies on the most common target of genome manipulation, the mouse, are needed. Surprisingly few studies describe

GAML in the mouse. Moreover, to my knowledge, no study of AAS alleviation of GAML in mice has been published. While some parallels with rat studies may help, higher resistance to Dexa in mouse suggests there must be differences. This work sets out to determine whether AAS can alleviate GAML in mice. Upon identifying the conditions that lead to GAML and to its alleviation by AAS, the focus will switch to understanding of the molecular mechanisms underlying them. Based on the published evidence available at the initiation of this project, I set forth the following hypotheses:

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**1. Testosterone alleviates dexamethasone-induced muscle atrophy in mice.** GAML has been frequently reported in rats and infrequently in mice. A series of studies describe reductions in CSA of mouse myofiber upon Dexa treatment. On the other hand, Testo has been used to reverse loss of muscle in mice and other species. Based on the information published prior to this work, there was a strong probability for Testo to prevent GC-induced loss of mouse muscle.

## 2. Testosterone’s myoprotective action is based on inhibiting dexamethasone’s proteolytic effects of dexamethasone. Studies of

GAML suggest that it involved a strong upregulation of proteolysis, aggravated by a decrease in protein synthesis. Protein degradation mechanisms upregulated by Dexa are diverse, including the proteasome, the autophagosome, and soluble proteases. Microarray studies showed that Testo reverses GC-induced changes in markers of each of the three proteolytic pathways and of protein translation. Given that the amplest changes induced by Dexa are on the proteasome ubiquitin system, there was a good probability that specific reversal of GAML by Testo will include proteasome’s repression.

1. **IGF-I upregulation facilitates testosterone’s myoprotectiveaction in glucocorticoid-induced loss of muscle mass.** The few studies on GAML reversal by AAS revealed that Testo action alters mTOR and Akt signaling. Based on the microarray studies which showed that Testo reverses changes in IGF-I and IGFBP induced by Dexa, and on the castration reversal studies indicating that AAS replacement causes IGF-I upregulation, there was a sizeable likelihood that the myoprotective action of Testo involved the modulation of IGF-I signaling.
2. **METHODS**

## Ethical considerations

All animal procedures have been described in protocols drafted by the author. The protocols were written in advance, submitted with Dr. Carlo Serra as principal investigator, and approved by the Institutional Animal Care and Use

Committee at the Boston University School of Medicine and Harvard Medical Area Standing Committee on Animals.

## Animal studies

Male, 6-8 week old (young adult), C57Bl/6J mice were purchased from The Jackson Laboratories (Bar Harbor, Maine). Mice were acclimated for 3 d to 7 d between delivery and initiation of the experimental interventions. Before and during the experiments, mice were maintained in a temperature-controlled facility, at 21 ◦C, with 12 h light / 12 h dark cycles. Mice were offered water and chow (Purina, Richmond, Indiana) ad libitum.

Every morning, for 1-7 days, mice were weighted, then injected subcutaneously every morning, between 9 and 11 AM, 200 µL corn-oil based solution, including 14 µL grade ethanol, which delivered either (A) 0*.*7 mg Testo propionate (T), or (B) 0*.*25 mg Dexa (D), or (C) both T and Dexa in the above doses (DT), or (D) neither drug. This latter group will be designated Vehicle (V).

Testo propionate and Dexa were from Sigma-Aldrich (St. Louis, Missouri).

Research-grade corn oil was purchased from MP Biomedicals (Solon, Ohio).

Unless specified, chemical reagents used in this work were from Fisher

(Pittsburgh, Pennsylvania), including pharmaceutical-grade ethanol used here. Before the first injection and 24 h after the last injection, mice lean and fat

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body mass was measured by nuclear magnetic resonance (NMR), using a

Echo-MRI whole body composition analyzer (Echo Medical Systems, Houston, Texas). In this procedure, mice are restrained inside a transparent plexiglass tube for 90 s, without sedation or anesthesia.

Mice were euthanized humanely by Euthasol® (pentobarbital sodium and phenytoin sodium solution; 200 mg/kg pentobarbital; Diamond Animal Health, Des Moines, Iowa) intraperitoneal, followed by quick cervical dislocation. Blood was collected immediately after death through thoracotomy and cardiac puncture, incubated 15 min at room temperature, centrifuged 15 min, at 10,000 g, 4 ◦C. Levator ani, gastrocnemius, tibialis anterior, quadriceps, and triceps brachii muscles were collected, weighted in wet state, flash-frozen by submersion in liquid nitrogen, and stored at −80 ◦C.

Muscles were crushed under liquid nitrogen, using a mortar and pestle pre-chilled in liquid nitrogen, and muscle powder was stored at −80 ◦C. Small quantities of muscle powder (25 µg to 30 µg) were lyzed for enzymatic activity assays, immunoblot, or quantitative real-time polymerase chain reaction (qRT-PCR), as described in dedicated sections.

## Enzymatic assays

Chymotrypsin-like proteasome enzymatic activity was measured using the 20S Proteasome Activity Assay kit (Chemicon International, Temecula, California). For each animal and muscle, 25 mg powdered muscle was extracted with 8 mL/g tissue of lysis buffer containing 150 mM sodium chloride, 50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonate HEPES (pH 7.4), 5 mM sodium ethylenediaminetetraacetate (EDTA), and 10 ‰ Triton X-100, by nutation at 4 ◦C for 30 min, with vortexing every 10 min. The extract was clarified by

centrifugation at 10,000 g, 10 min, 4 ◦C . The clarified extract was collected and stored at −20 ◦C for 1 h, while its total protein concentration was determined using the bicinchoninic acid (BCA) method (detailed in the immunoblot section). Next, the clarified extract were diluted to match the lowest total protein concentration with lysis buffer. Per kit manufacturer instructions, 80 µL clarified extract (depending on muscle, 300 µg to 600 µg total protein) were mixed, in 96-well plate wells, with 10 µL proprietary assay buffer and 10 µL fluorogenic proteasome substrate, Succynyl-Leu-Leu-Val-Tyr-7-amino-4-methylcoumarin, 500 µM stock. This was done in duplicates for each sample, and for a negative control (“background”), containing lysis buffer instead of muscle extract. The plate was sealed and incubated at 37 ◦C for 30 min. Next, fluorescence was measured with a Saphire multi-well plate reader (Tecan, Männedorf, Switzerland) with excitation at 380 nm and emission at 460 nm. Background fluorescence, including substrate, was subtracted from sample measurements. Preliminary tests with manufacturer’s positive control and muscle lysate indicated that 30 min fluorescence measured with optimal gain and with background subtraction is proportional with the rate of fluorescence change and with the amount of measured enzyme. Thus, more detailed kinetics were not needed.

Lysosome enzymatic activity was measured in a similar protocol, using

Cathepsin L Activity Fluorometric Assay kit (Abcam plc, Cambridge, England).

For each animal and muscle, 25 mg powdered muscle was extracted with 8 mL/g

tissue of proprietary lysis buffer, clarified, stored, and assayed for total protein, in the same manner as for the proteasome activity assay. Extracts were diluted to the lowest total protein concentration, using the same lysis buffer that was used to extract them. Per kit manufacturer instructions, 48 µL clarified extract

(depending on muscle, 100 µg to 300 µg total protein) were mixed, in 96-well

plate wells, with 50 µL proprietary assay buffer and 2 µL fluorogenic cathepsin substrate, FR-amino-4-trifluoromethyl coumarin, 10 mM stock. This was done in duplicates for each sample, and for a negative control (“background”), containing lysis buffer instead of muscle extract. The plate was sealed and incubated at 37 ◦C for 30 min. Fluorescence was measured with excitation at 400 nm and emission at 505 nm. Background fluorescence, including substrate, was subtracted from sample measurements.

Calpain enzymatic activity was measured in a similar protocol, using Cathepsin L Activity Fluorometric Assay kit (Promega, Madison, Wisconsin). For each animal and muscle, 25 mg powdered muscle was extracted with 8 mL/g tissue of lysis buffer containing 150 mM sodium chloride, 10 mM HEPES pH 7.4, 10 mM dithiothreitol (DTT), 1 mM EDTA, and 10 ‰ Triton X-100, as described for proteasome activity assay. Extracts were clarified, and stored, as described for the proteasome activity assay. Total protein was measured by Bradford assay.

Briefly, 1 µL sample was mixed with 300 µL Coomasie Plus (Life Technologies, Carlsbad, California) in duplicate wells of a 96-well plate, incubated 10 min at room temperature, and measured spectrophotometrically for absorption at 595 nm. Extracts were diluted to the lowest total protein concentration, using the same lysis buffer that was used to extract them. Per kit manufacturer instructions, 50 µL clarified extract (depending on muscle, 100 µg to 300 µg total protein) were mixed, in 96-well plate wells, with 50 µL proprietary assay buffer, containing lyophilized luciferase, ATP, 80 µM pro-luminescent calpain substrate, Suc-Leu-Leu-Val-Tyr-aminoluciferase, and 8 µM calcium chloride. This was done in duplicates for each sample, and for a negative control (“background”), containing lysis buffer instead of muscle extract. After 10 min of incubation at room temperature, steady state is reached, with a constant rate of free aminoluciferase production, which is immediately converted to free luciferin by the excess luciferase. Luminescence was measured with a GeniosPro multi-well plate reader (Tecan, Männedorf, Switzerland) set at to integrate signal over 100 ms. Background luminescence, including substrate, was subtracted from sample measurements.

## Immunoblot

For each animal and muscle, 25 mg powdered muscle was extracted with

4 mL/g tissue of radioimmunoprecipitation assay (RIPA) buffer, containing

150 mM sodium chloride, 20 mM Tris pH 7.5, 100 ‰ v/v glycerol, 10 ‰ v/v

NP-40, 10 g/L sodium deoxycholate, 1 g/L sodium dodecyl sulfate (SDS), 1 mM EDTA, 1 mM sodium ethyleneglycol tetraacetate, 10 mg/L leupeptin, 10 mg/L pepstatin, 10 mg/L aprotinin, 1 mM phenylmethanesulfonylfluoride, 1 mM sodium orthovanadate, 5 mM sodium fluoride. The mixture was nutated at 4 ◦C for 30 min, with vortexing every 10 min. The suspensions are clarified by centrifugation at 12,000g, 15 min , at 4 ◦C. The supernatant is collected, aliquoted, and stored for months at −80 ◦C.

Total protein content was measured with a BCA proprietary kit (Thermo Scientific Pierce, Rockford, Illinois). First, a 50:1 combination of reagents VW A and B was prepared. Next, 200 µL VW reagent mix was combined with 2 µL unknown solution in a 96-well plate well. This was done in duplicates for each sample, and for a series of standards of 0*.*5 g/L to 10 g/L bovine serum albumin (BSA) prepared with the same lysis buffer as the unknowns. The plate was incubated 30 min at 37 ◦C. Light absorbance was measured with a Saphire multi-well plate reader (Tecan, Männedorf, Switzerland) with absorption at

562 nm. The standard curve was fitted to a quadratic equation. Typically extracts were diluted to the lowest total protein concentration, using the same lysis buffer that was used to extract them.

The lysates were resolved by polyacrylamide gel electrophoresis (PAGE). The gels were cast in the morning of the electrophoresis. The stacking section was a 5 g/dL acrylamide gel, with 125 mM Tris pH 6.8. The separating section has a variable concentration of acrylamide (12 g/dL for low molecular weight unknowns, 8 g/dL for others), and 400 mM Tris pH 8.8. Both sections included 1 g/L SDS, and polymerized by the addition of 1 g/L ammonium persulfate and 0*.*4 ‰ v/v tetramethylethylenediamine. In order to settle and migrate, lysates were mixed 1:1 with PAGE sample buffer, which contained 30 mM Tris pH 6.8,

10 g/L SDS, 100 ‰ v/v glycerol, 50 mg/L bromophenol blue, and 350 mM DTT. The mixture of sample and sample buffer was incubated for 5 min at 95 ◦C, then loaded on the gel, with equal volumes and masses of total protein in each well.

Electrophoresis was performed in a Miniprotean Tetra cell (Bio-rad, Hercules, California), at constant 80 volts, using an electrode buffer with 14*.*4 g/L glycine, 3 g/L Tris base, 10 g/L SDS.

From the gel, resolved proteins were transfered to nitrocellulose membranes using a Bio-rad Mini-trans blot cassette, at 8 V/cm constant overnight at 4 ◦C. The Towbin transfer buffer had 200‰ v:v methanol, 25 mM Tris pH 8.3, 192 mM

glycine.

Quality of transfer was assessed by temporary Ponceau staining (20 g/L Ponceau S in 300 g/L trichloroacetic acid and 300 g/L sulfosalicylic acid). At this time, using indications from the molecular weight standards, the membrane was sectioned, thus allowing probing with multiple antibodies.

Ponceau was removed by washing twice, with shaking 5 minutes at room temperature, in TBST (130 mM sodium chloride, 20 mM Tris pH 7.5, 1‰ v/v Tween-20). The membranes were blocked by shaking for one hour in blocking solution (TBST with 5% fat-free instant milk) at room temperature. Next, membranes were probed by shaking overnight at 4 ◦C in TBST with 50 g/L BSA and primary antibodies of choice (listed later). Next day, the membranes were washed in TBST, twice briefly and three times with 5 minutes shaking at room temperature. The washed membranes were then probed by shaking at room temperature for an hour in a solution with TBST, 5% fat-free instant milk, and the secondary antibody of choice (listed later). The membranes were washed as before.

Next, the membranes were probed with proprietary Amersham ECL Western Blotting Detection Reagent (GE Healthcare, Wilmington, Massachusetts). Briefly, the two reagent components were warmed to room temperature, mixed 1:1, and layered on the nitrocellulose membrane. The membrane was incubated for 1 minute, then excess liquid was drained. The membrane was placed in the film cassette. In a dark room, under red light, a X-Omat photographic film (Kodak, Rochester, New York) was placed on the nitrocellulose membrane. Exposure was between 10 seconds and 5 minutes, depending on the strength of the signal. After developing the first image, longer or shorter exposures were used, as needed.

Photographic films were digitized using a FluorChem-SP Imaging System (Alpha Innotech, San Leandro, California), using automatic exposure. Band densitometry was performed with the Image Studio Lite application (Li-Cor,

Lincoln, Nebraska).

At times, antibodies were stripped by incubation with shaking for 45 minutes at 50 ◦C in a tight container with pre-warmed stripping buffer (20 g/L SDS, 60 mM Tris pH 6.8, 100 mM beta-mercaptoethanol). Stripped membranes were rinse with water, washed with TBST, re-blocked, and re-probed.

Primary antibodies used were: anti-myosin heavy chain (in-house, from

MF20 hybridoma cells), anti-GAPDH (Pierce), anti-calpain 1 (Abcam), anti-eIF3f (Rockland), anti-LC3, anti-phospho-eIF2, anti-4EBP, anti-phospho-4EBP, anti-ATF4, anti-cathepsin L, anti-calpastatin, anti-IGF-1R, anti-phospho-IGF1-R, anti-Akt, anti-phospho-Ser473 Akt (all from Cell Signaling Technologies). Anti-mouse and anti-rabbit secondary antibodies were from Cell Signaling Technologies.

## Quantitative real-time polymerase chain reaction

Total RNA was purified by a combination of phenol-chlorophorm fractionation and spin column chromatography. For each animal and muscle, 25 mg powdered muscle was homogenized briefly in 1 mL Trizol (Life Technologies). The suspension was incubated 5 minutes at room temperature, then supplemented with 200 µL chloroform. The tube was shaken vigorously for 15 seconds, then incubated 3 minutes at room temperature. Phase separation was induced by centrifugation, 15 minutes, 12,000 g, at 4 ◦C. The top, aqueous fraction (500 µL) was collected, then passed through a gDNA exclusion column, by centrifugation for 30 seconds, 8,000 g. The recovered solution was supplemented 1:1 with 700‰ v/v ethanol, then passed through an RNeasy column, by centrifugation, 10 seconds at 8,000 g. The column was washed with 700 µL of buffer RW1, and twice with 500 µL of buffer RPE, each time by centrifugation, 15 seconds at 8,000 g. The column was dried by centrifugation 2 minutes at 8,000 g. RNA was eluted with 40 µL water, by centrifugation, 1 minute at 8,000 g.

Total RNA concentration was measured with an ND-1000 spectrophotometer (Nanodrop, Wilmington, Delaware) at 260 nm, using its Windows proprietary

application.

To obtain cDNA using Accuscript 1st Strand cDNA synthesis kit, 320 ng total RNA was combined with 300 ng random primers, 20 nmol of each triphosphate nucleotide, and the proprietary AccuScript RT Buffer 10×. After incubation 5 minutes at 65 ◦C, and cooling towards room temperature another 5 minutes, the RNA mix was supplemented with 2 nmol DTT, 20 units RNase Block ribonuclease inhibitor, and 1 μL reverse transcriptase (“AccuScript RT”) proprietary stock. For negative control purposes, the same mixture was prepared, while withholding the RT Buffer and the RT stock. This no-RT tube provided a measurement of the leaked DNA, which, in the course of this work, was negligible. Reverse transcription was performed by incubating 10 minutes at 25 ◦C, then 60 minutes at 42 ◦C. Reverse transcription was terminated by incubating 15 minutes at 70 ◦C.

The product was stored at 20 ◦C between measurements.

Specific cDNA amounts were measured with an 7500 Fast Real-Time PCR cycler (Applied Biosystems, Foster City, California), using 96-well plates. Each unknown was measured in duplicates. Each well contained 1 µL cDNA stock, 10 pmol each sense and anti-sense primer (listed later), and proprietary SybrGreen

2X. The qRT-PCR cycler incubates samples 2 minutes at 50 ◦C, then 10 minutes at

95 ◦C. Then, 40 cycles are run, comprising 15 seconds at 95 ◦C and one minute at 60 ◦C, with measurements of DNA after each cycle. The amount of specific cDNA is estimated from the number of cycles (Ct; cycle threshold) required to reach a DNA concentration threshold, automatically chosen by the Applied Biosystems software at midway between the lag phase and the end plateau DNA concentrations. Moreover, the Applied Biosystems software is able to interpolate the number of cycles, thus providing, for each well, a non-integer number of cycles that will yield the threshold DNA concentration.

Duplicates were averaged for each unknown. Next, from the cycle count

|  |  |
| --- | --- |
| Gene | Primers |
| Trim63 | Forward: TCTCTCAGCTGGAGGACTCC Reverse: CTGTACTGGAGGATCAGAGC |
| Fbxo32 | Forward: GGGAAGCTTTCAACAGATTGG Reverse: AATGTTCATGAAGTTCTTTTGGG |
| Igf1 (IGF-IEa) | Forward: GCTTGCTCACCTTTACCAGC Reverse:AAATGTACTTCCTTCTGGGTCT |
| Foxo1 | Forward: CACCCTGTCGCAGATCTAG Reverse: CGAATAAACTTGCTGTGAAGG |
| Foxo3a | Forward: GACAAACGGCTCACTTTGCC Reverse: GACAGGTTGTTGTGCCGGATGG |
| Foxo4 | Forward: GCCCTACTTTCAAGGACAAGG Reverse: CATCCACCAGAGCTCTTC |
| Ddit4 | Forward: CTGCGAGTCCCCTGGACAGC Reverse: CACTGAGCAGCTCAAAGTCG |
| Igf1r | Forward: GGGCTGACTGGTGGATGC Reverse: CGCTGGGCACGGATAGAGC |
| Becn1 | Forward: CCACAGCCCAGGCGAAACC Reverse: GATGAATCTTCGAGAGACACC |
| Map1lc3b | Forward: GGAAGATGTCCGGCTATCC Reverse: CTCATGTTCACGTGGTCAGG |
| Ctsl | Forward: AACCGCTGATGCTTAAGATCC Reverse: CACTCAGTGAGATCAGTTTGC |
| Klf15 | Forward: ACAGGCGAGAAGCCCTTGC Reverse: CTCGCACACGGGACACTGG |
| Gapdh | Forward: GCTCACTGGCATGGCCTTCCG Reverse: GTAGGCCATGAGGTCCACA |

Table 2: List of RT-PCR primers.

required to reach the threshold DNA concentration for the specific cDNA of interest, the cycle count for a housekeeping gene was subtracted, thus providing a relative measurement of expression for the gene of interest within the transcriptome of that animal. Statistics were computed on these differences, also known as ΔCt. To enable comparisons, these ΔCt measurements were further re-based, by subtracting from each the average measurement in the V group.

(This double-rebasing method is also referred to as the ’ΔΔCt’.)

Primers used are listed in table 2.

## Cell culture studies

C2C12 cells were obtained from ATCC (Manassas, Virginia). They were grown on solid substrate, on cell culture coated plates, in atmospheric air, supplemented to 5% CO2, at 37 ◦C. During proliferation, cells were maintained under

500 µL/cm2 growth medium, that is, Dulbecco’s Modified Eagle Medium (DMEM; VWR International, Radnor, Pennsylvania), supplemented with 100‰ v/v fetal bovine serum (FBS), 100 units/mL penicillin, and 100 mg/L streptomycin, changed every other day. DMEM contained 400 µM phenylalanine, 4*.*5 g/L glucose, 4 mM glutamine, and lacked pyruvate.

Confluence and multiple passages reduce their ability to proliferate and differentiate. Therefore, cells were propagated in an undifferentiated state, by splitting 1:40 at confluence, about every 5 days. Early on, frozen stocks were made, at 1 million cells/mL in growth medium supplemented with 100‰ v/v dimethyl sulfoxide (DMSO), stored under liquid nitrogen. For detachment, cells were rinsed twice with pre-warmed 200 µL/cm2 phosphate buffered saline (PBS), then incubated at 37 ◦C for about 5 minutes with 50 µL/cm2 pre-warmed 0.05% trypsin / EDTA (Life Technologies). After detachment, cells were tested for viability with trypan blue, counted on a hemocytometer, and re-seeded at about 5,000 cells/cm2.

For experiments, cells were differentiated into myotubes by seeding at higher density (about 5,000 cells/cm2). Next day, when cells were nearly confluent, the medium was changed to differentiation medium, that is, DMEM supplemented with 2% horse serum (HS), which was charcoal / dextran stripped of hormones (Gemini Bio-Products). After two days in differentiation medium, myotubes form, while myoblast become quiescent. The medium was then changed every day for five more days, leading to nearly complete elimination of undifferentiated and mononuclear cells.

Dexa and Testo were delivered from 400X stocks in ethanol, at various concentrations, as indicated at each experiment in the results section.

Protein synthesis was inhibited with 100 µM cycloheximide from 360X water-based stock, or 5 µM puromycin dihydrochloride from 500X water-based stock. Proteasomal activity was inhibited in cell culture with 5 µM MG132 from a

80X working stock in differentiation medium, which in turn is derived from a 8,000X DMSO-based stock. Lysosomal activity was inhibited with 25 µM chloroquine from a 2,000X water-based stock. Calpain and cathepsin activity was inhibited by 10 µM E-64, from a 100X water-based stock. IGF-1R was inhibited by 50 nM picropodophyllin (PPP), from a 200X working stock in differentiation medium, which in turn is derived from a 1,000X DMSO-based stock.

## Immunofluorescence microscopy

After the experimental treatments, the adherent cells were washed twice, on ice, with ice-cold PBS. The cells were covered with cold 1:1 v:v ethanol:acetone, and incubated at 20 ◦C for 20 minutes. This process both fixated and permeabilized the cells. The liquid was removed, and the cells were rinsed with PBS.

On the first day of staining, PBS was removed, and cells were washed briefly twice with PBS. For blocking purposes, cells were incubated for 40 minutes at room temperature under 1% BSA in PBS. The blocking liquid was then replaced with the primary antibody solution, that is, the same blocking solution, supplemented with 1:20 hybridoma cell culture medium containing the anti-myosin heavy chain antibody MF-20. Cells were probed with the primary antibody overnight at 4 ◦C.

Next day, the cells were incubated twice 5 minutes each with the blocking solution, to wash unbound primary antibody. Cells were then probed with the secondary antibody, that is, blocking solution supplemented with 1:500 anti-mouse antibody conjugated with rhodamine, and with 1 mg/L DAPI. The cells were incubated 1h at room temperature in the dark, then briefly washed 4 times with PBS. Cells covered in PBS were store at 4 ◦C or observed with an Eclipse TE2000-E fluorescence microscope (Nikon Instruments, Melville, New York).

Cell contours were obtained using Magnetic Lasso in Adobe Photoshop CS4. All the cell outlines that could be recognized by Photoshop from two randomly chosen fields were pasted in another file, that was read in Matlab 8. An automated script determined multiple estimates of the diameter, and calculated an average diameter for each fiber.

## Measurement of muscle protein synthesis and degradation

Cells were labeled by incubation in differentiation medium supplemented with 0*.*2 mCi/L radioactive (³H-[ring]) phenylalanine (126 Ci/mmol126 Ci/mmol; PerkinElmer, Shelton, Connecticut), starting at the times and for the durations specified at each experiment in the results section. For removal of non-specific tracer at the end of labeling stage, cells were rinsed with PBS twice, then incubated with 2 mm PBS for 15 min at 37 ◦C, 5% CO2.

For chase experiments, PBS was then replaced with non-radioactive differentiation medium, for the durations specified at each experiment. Chase experiments ended with a double rinse with PBS. Preliminary studies revealed that supplementation of differentiation medium with 10 mM phenylalanine made no difference to the rate of tracer release, compared to differentiation medium alone (data not shown). This is plausible, because differentiation medium already contains 0*.*4 mM phenylalanine, and suggested that additional osmotic stress could be avoided at no cost to sensitivity of the method.

In all cases, cells and media were collected separately, and each was fractionated with trichloroacetic acid (TCA). Medium was removed from the plate, then supplemented with 6 M TCA, to a final 0*.*6 M TCA. Cells were covered with 0*.*6 M TCA. Both media and cells were then incubated overnight at −20 ◦C. Next day, cells and media were thawed at 4 ◦C over two hours. Cells were scraped. The cell culture wells were further scraped and rinsed with 0*.*6 M TCA. Protein suspensions were centrifuged at 15,000 RCF, for 15 min, at 4 ◦C. Pellets were washed with 0*.*6 M TCA, by resuspension, incubation at 4 ◦C for 15 min, and centrifugation as above. Next, pellets were washed once more with acetone, by the same procedure. All two TCA-based supernatants and the acetone supernatant were pooled. Pellets were dissolved in 1 M NaOH / 1‰ v/v Triton, with sonication 15 min at 50 ◦C, followed by incubation 15 min at 50 ◦C.

From the supernatant or from the resuspended pellet, 100 µL was mixed with 5 mL Econo-lite scintillation fluid, then counted in a TriCarb 3100TR liquid scintillation counter (PerkinElmer) for 3 minutes per sample. Counts per minute were converted to disintegrations per minute using a standard curve generated using 1 µCi/L to 80 µCi/L labeled phenylalanine dissolved in the same solvents as the samples (2:1 TCA:acetone, or 1 M NaOH / 1‰ v/v Triton).

From the same protein pellet suspension, aliquots were used to measure total protein concentration by BCA method, as described in the immunoblot section, but with standards prepared in the same solvent as the unknowns.

## Statistics

Statistics and plots were generated in R 3.2 with ggplot2, knitr, and Lyx[628]. Results are presented as means ± standard error of the mean. In plots, points represent means, while error bars are standard error of the mean. In both cases, error bars include only biological variability. Experiments with more than two groups were analyzed by Kruskal-Wallis test for overall non-parametric analysis of variance. Groups were compared in pairs using post-hoc Dunn’s test with Bonferroni correction for multiple comparisons. In the cases where data passed the Shapiro-Wilk test of normality at significance level of 0.05, parametric tests were used instead, including analysis of variance (ANOVA) and post-hoc Tukey’s honest significant difference (HSD) test.

In a plot, means with the same letter are not significantly different from each other. Tests were considered statistically significant when the null hypothesis could not be rejected at any level of significance greater than or equal to 0.05.

## Literature review

The introduction section was based on review of all literature indexed by

PubMed. Search expressions included ‘testosterone OR androgens‘, ‘dexamethasone OR betamethasone OR triamcinolone OR prednisone OR prednisolone OR hydrocortisone OR cortisone OR triamcinolone OR fludrocortisone‘, ‘Cushing‘, ‘ribosome OR polysome OR lysosome OR autophagosome OR proteasome OR ligase OR cathepsin OR FOXO OR IGF1 OR calpain OR mTOR OR AMPK OR Akt‘and combinations thereof. Relevant primary data were summarized.

Literature plots were digitized with WebPlotDigitizer, a web application

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created by Ankit Rohatgi[629].

**5. IN VIVO EXPERIMENTS**

## Testosterone reverses dexamethasone-induced muscle atrophy

No published study on mice describes a dose of Dexa that is proven to induce muscle atrophy. I attempted to repeat the daily injections protocol described by Jones as effective on rats[521] (Fig.2 ).

Mice treated with the Dexa doses that were reported to induce muscle atrophy in rats did not exhibit significant muscle loss based on muscle dissection and NMR (data not shown). Higher doses of Dexa were used. Effect size on wet muscle weight became stronger than experimental error at a dose of 10 mg/(kg d) Dexa. In contrast, Testo doses that could reverse GAML were found to be similar to those effective in rats (28 mg/(kg d)).

Upon finding the Dexa dose effective in inducing muscle atrophy, I analyzed the body weight at sacrifice. There was no difference between groups in terms of absolute weight, due to to the large variability of the initial body weight (Kruskal-Wallis p = 0.777) (Fig. 3, right). The difference became apparent when percent change in body weight was compared(Fig. 4, top, last time point). In this case, the three treatments (vehicle, Dexa, or the combination Dexa + Testo) were significantly different (Kruskal-Wallis p = 0.00385). Instead of the expected unchanged body weight over the seven days of treatment with vehicle alone, there was a trend for growth, with an apparent gain of 0.212 grams every day during the treatment. A similar trend has been seen in the other repetitions of the experiment, as well as in other published studies on young rodents. A sizable contribution to this growth was brought by the 200 µL vehicle injected every day. Once this artifact is taken into account, the gain of a negligible 0.0333 grams per day in the Dexa-treated group is in fact indicative of an actual massive loss of body

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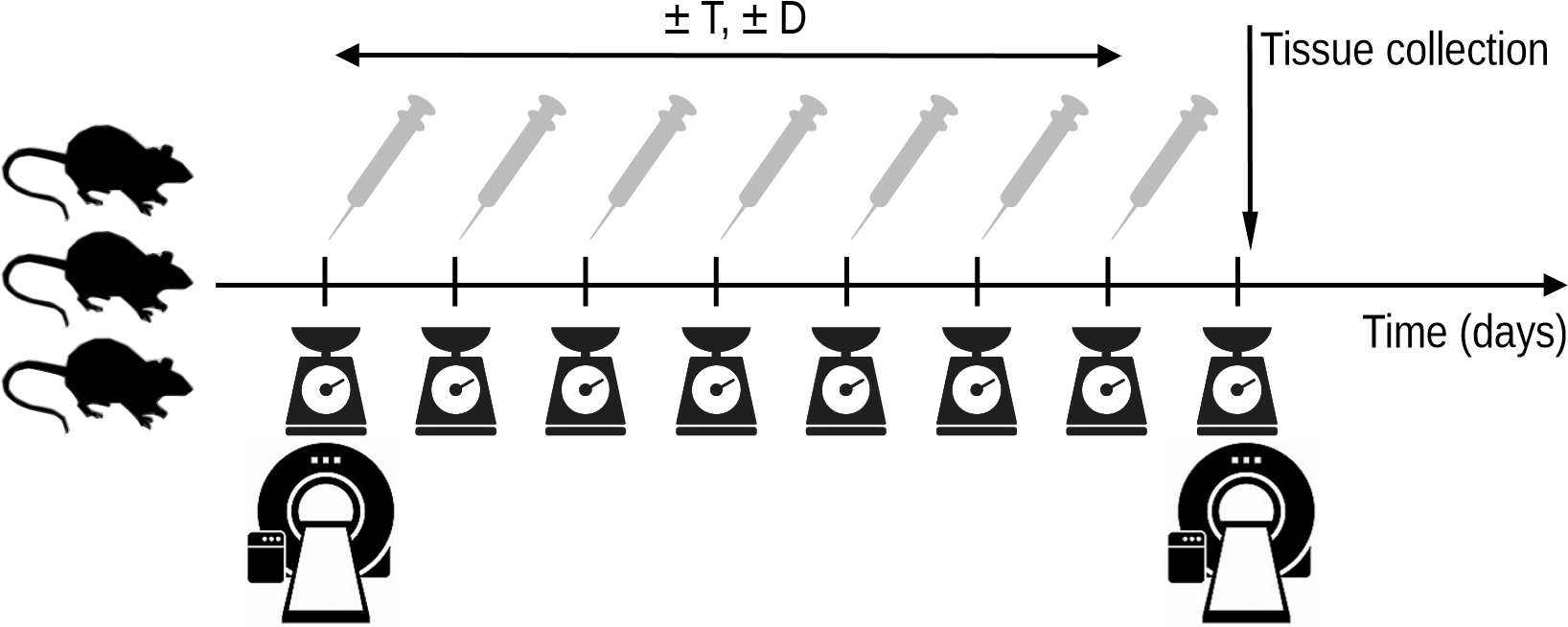


Figure 2: Experimental protocol for the in vivo studies.

a

b

a,b

one day

0

10

20

30

Dexa Comb

Veh

body weight (g)

a,b

a

b

three days

0

10

20

30

body weight (g)

a

a

a

seven days

0

10

20

30

body weight (g)

Figure 3: Body weight at sacrifice, following 10 mg/(kg d) Dexa and / or 28 mg/(kg d) Testo treatments. Treatments designated by the same letter do not differ significantly from each other (n=5-6).

†

✶

†

†

†

†

†

†

✶

-5

0

5

10

1

2

3

4

5

6

7

8

day

body weight gain (% of pre-treatment)

-5

0

5

10

1234

day

body weight gain (% of pre-treatment)

-5

0

5

10

12

day

body weight gain (% of pre-treatment)

Veh

Dexa

Comb

Figure 4: Time course of body weight Dexa and / or Testo treatments (n=5-6). Stars indicate statistically significant differences between Veh and Dexa (p <0.05). Daggers indicate statistically significant differences between Dexa and Comb (p <0.05).

weight.

Seven-day weight gain in Dexa-treated group (1.05 ± 1.14% of initial body weight) was significantly smaller than that of the vehicle-treated group (6.12 ± 0.879%; Dunn’s test p = 0.0447). Conversely, co-administration of Testo, hereafter and in plots abbreviated Comb, brought back the body weight gain over seven days to levels similar to those in vehicle-treated animals (8.62 ± 0.674%; Dunn’s test vs. Dexa alone, p = 0.00167).

The time course of body weight changes suggested that both drugs’ action had

a rapid onset (Fig. 4; Kruskal-Wallis for first day percent change in body weight, p = 0.0239). Specifically, mice receiving vehicle alone gained 2.28 ± 0.485% body weight in the 24 hours, due to accretion of nonresorbable vehicle. In contrast, mice receiving Dexa lost 1.86 ± 0.485% body weight (Dunn’s test p = 0.0387).

Mice receiving a combination of Dexa and Testo were essentially indistinguishable from those receiving vehicle alone, having gained 2.61 ± 0.767% body weight. This demonstrated an advantage of the combination treatment over Dexa alone

(Dunn’s test p = 0.0213).

Based on the whole time course, I hypothesized that body weight changes and muscle atrophy occur in a gradual manner, with significant metabolic and molecular changes preceding the seven-day end of experiment. Accordingly, I repeated the above experiment on different cohorts of mice, which were sacrificed after only 1 or 3 days of treatment. Once more, absolute changes in body weight could not be correlated with the muscle changes (Fig. 3, left and middle). When normalized, relative changes in body weight for these groups of mice were less ample than those seen with 7 days of treatment, to the point that most parameters (body weight, lean body mass, individual muscles mass) were not changed in a statistically significant manner (Fig. 4, bottom). These shorter treatments provided insight into the molecular development of atrophy, but, because they were run as independent experiments, they will not be analyzed in conjunction. Because this work is focused on longer-term effects of Dexa, most of the reported data in the remainder of the section will be from mice treated for 7 days with Dexa and / or Testo, with exceptions and subacute effects emphasized.

Body composition analysis indicated that all animals included in these experiments progressively lost total water. There was no difference between water loss in the three experimental groups (Fig. 5, top; Kruskal-Wallis for seven day mass of water lost, p = 0.302). This uniform loss of water negates a scenario in which the observations could be ascribed to increased water retention due to non-specific action of either steroid with MR. In all cases, the losses of total water track the loss of lean body mass, indicating that muscle atrophy, rather than renal dysfunction, underlies the loss of water. Fat mass analysis indicated that in the vehicle-alone treated group, the rate of apparent fat gain was essentially equal with the mass of injected vehicle accrued over the seven days. This demonstrates that experiments manipulations, in the absence of pharmacological treatments, have no effect on lipid metabolism. Dexa-treated mice accrued an additional 0.73 g fat, compared to vehicle (Dunn’s test, p = 0.0681. Similarly, mice treated with the combination accrued 0.922 g additional fat, compared to vehicle (Dunn’s test, p = 0.0172). Therefore, Testo co-administration had no effect on lipid accretion (Dunn’s test, Dexa vs. combination, p = 0.802).

Ampler changes were induced by the two drugs on lean body mass (Fig. 5, bottom; Kruskal-Wallis for seven day lost lean mass, p = 0.000811). Vehicle alone has no effect on lean body mass (0.327 ± 0.203 grams lost over seven days, that is, 1.65 of the total lean body mass). Dexa alone induce a massive loss of lean body mass (3.21 ± 0.166 grams lost over seven days; Dunn’s test p = 0.000242). When Testo was co-administered, the loss of lean body mass persisted, but there was a trend towards a lower loss rate (1.57 ± 0.133 grams lost over seven days; Dunn’s test vs Dexa, p = 0.108).

The loss of lean body mass induced by Dexa was apparent even in the 3-day experiment (1.45 ± 0.152 grams lost over three days; Dunn’s test vs. vehicle, p = 0.00961). Interestingly, in that experiment, Testo protective effect was very small and not statistically significant (1.28 ± 0.369 grams lost over three days; Dunn’s test vs. Dexa, p = 0.675).

a

a

a

one day

0

1

2

3

4

5

water loss (g)

a

a

a

three days

0

1

2

3

4

5

water loss (g)

a

a

a

seven days

0

1

2

3

4

5

water loss (g)

a

a

a

0

1

2

3

4

fat mass gain (g)

a

a,b

b

0

1

2

3

4

fat mass gain (g)

a

a,b

b

0

1

2

3

4

fat mass gain (g)

a

a

a

0

1

2

3

4

Veh

Dexa

Comb

lean mass loss (g)

a

b

a,b

0

1

2

3

4

Veh

Dexa

Comb

lean mass loss (g)

a

b

a,b

0

1

2

3

4

Veh

Dexa

Comb

lean mass loss (g)

Figure 5: Changes in water, lean, and fat body mass, after Dexa and / or Testo treatments (n=5-6).

Dissection of individual muscles confirmed that Dexa achieved widespread muscle atrophy (Fig. 6). Despite the small sample size (n=5-6), the atrophying effect of Dexa became statistically significant at day 3 on gastrocnemius (Dunn’s test, p = 0.0369). At day 7, statistical significance is also achieved in triceps brachii (Dunn’s test, p = 0.0209), quadriceps (Dunn’s test, p = 0.00205), and levator ani (Dunn’s test, p = 0.0245). In terms of amplitudes, the five measured muscles ranged from extremely responsive, such as quadriceps (22.7% muscle weight loss), triceps (18% muscle weight loss) and gastrocnemius (16.5% muscle weight loss), to the refractory tibialis anterior (6.2% muscle weight loss). For each muscle and time point, the average muscle weight in the Dexa group was smaller than the average weight of the controls.

Upon Testo co-administration, four out of five muscles measured were exhibited a trend towards restoration to their basal weight (Fig. 6). The amplitude of restorative response to Testo co-administration was strongest in quadriceps

(15.2% restoration in muscle weight; Dunn’s test, p = 0.27), followed by levator

(10.2% restoration in muscle weight; Dunn’s test, p = 0.134), and gastrocnemius (9.07% restoration in muscle weight; Dunn’s test, p = 0.199) appear the most responsive.

At the seventh day, Testo co-administration led to an additional 3.12% loss in tibialis weight (Dunn’s test, p = 0.523). A similar trend indicating Testo’s inability to rescue tibialis mass is also present in the three-day data (2.63% loss; Dunn’s test, p =0.975). The small amplitude and the complete absence of statistical significance suggest that observations describe lack of Testo effect, rather than a true atrophic effect. Because tibialis is not manifesting GAML at macroscopic level, molecular analysis will not include it.

In conclusion, 10 mg/(kg d) Dexa injection led to rapid onset of muscle

a

a

a

one day

0

25

50

75

100

levator ani (mg)

a

a

a

three days

0

25

50

75

100

levator ani (mg)

a

b

a,b

seven days

0

25

50

75

100

levator ani (mg)

a

a

a

0

50

100

150

200

quadriceps (mg)

a

a

a

0

50

100

150

200

quadriceps (mg)

a

b

a,b

0

50

100

150

200

quadriceps (mg)

a

a

a

0

50

100

150

gastrocnemius (mg)

a

b

a,b

0

50

100

150

gastrocnemius (mg)

a

b

a,b

0

50

100

150

gastrocnemius (mg)

a

a

a

0

50

100

150

triceps br. (mg)

a

a

a

0

50

100

150

triceps br. (mg)

a

b

b

0

50

100

150

triceps br. (mg)

a

a

a

0

20

40

60

Dexa Comb

Veh

tibialis ant. (mg)

a

a

a

0

20

40

60

Dexa Comb

Veh

tibialis ant. (mg)

a

a

a

0

20

40

60

Veh

Dexa Comb

tibialis ant. (mg)

Figure 6: Effects of Dexa and / or Testo treatments on the wet weight of levator ani, quadriceps, gastrocnemius, triceps brachii, and tibialis anterior muscles (normalized to body weight; n=5-6).

atrophy. Similar to humans with hypercortisolism, total body weight had limited use in assessing atrophy progression. Based on the time course of lean body mass and individual muscle changes, the atrophy develops throughout the first week, indicating that this murine model replicates chronic GC exposure. Some muscles are more responsive than others, with tibialis a notable refractory exception. In contrast, 28 mg/(kg d) Testo co-administration reduced the loss of muscle throughout the seven days course. While rat experiments have described a series of pathways by which Dexa can achieve muscle atrophy, it was unclear which of these, if any, is reversed by Testo co-administration.

**Testosterone reverses glucocorticoid-induced activation of the ubiquitin-proteasome system**

The rapid loss of muscle mass induced by Dexa in mice exceed the rate of protein turnover in normal adult rodent muscle (about 3% per day[630]). Therefore, Dexa’s action cannot rely exclusively on translational shutdown. I attempted to establish which protein degradative pathways could be mediating the atrophic effect of dexamethasone. Experiments on rats suggest that Dexa causes protein degradation by stimulating the proteasome-ubiquitin-related transcriptional program, whereas Testo represses it.

In normal circumstances, the rate-limiting factor in proteasome-mediated protein degradation is target protein conjugation with ubiquitin, catalyzed by E3 ligases. In mice treated with Dexa, the E3 ligase more commonly associated with loss of muscle mass, MuRF-1 (Fig. 7, top), was upregulated by Dexa at day 1 in gastrocnemius (1.9-fold amplification; Dunn’s test, p = 0.116) and in quadriceps (61.5-fold amplification; Dunn’s test, p = 0.00906). The effect was even more consistent at day 3, with MuRF-1 transcript in gastrocnemius being 7.58-fold amplified (Dunn’s test, p =0.0154), and in quadriceps 4.71-fold amplified (Dunn’s test, p =0.0726). The effect was insignificant statistically, and in amplitude, by the seventh day of Dexa administration.

Testo co-administration had an inhibitory effect on MuRF-1, with a later onset. No significant effect of Testo co-administration was seen in day 1. In day 3 samples, MuRF-1 was repressed by Testo in gastrocnemius (1.97-fold reduction; Dunn’s test, p =0.591) and in quadriceps (5.82-fold reduction; Dunn’s test, p =0.0268). Testo repression of MuRF-1 strengthened in the day 7 samples, with transcripts reduced in gastrocnemius (2.11 fold reduction; Dunn’s test, p = 0.116) and in quadriceps (2.13 fold reduction; Dunn’s test, p = 0.0748).

A similar pattern, although with lower intensity, was seen in MAFbx gene regulation by the two steroids. MAFbx was reliably upregulated by Dexa at day 1 in gastrocnemius (2.45 fold amplification; Dunn’s test, p = 0.175) and quadriceps

(50.1 fold amplification; Dunn’s test, p = 0.00906). At day 3, only the quadriceps

MAFbx response was still present to a significant degree (4.32 fold amplification; Dunn’s test, p = 0.113). The effect became insignificant statistically across muscle groups by the seventh day of Dexa administration.

Testo co-administration had no statistically significant effect on MAFbx at days 1 and 7. By day 3, MAFbx was repressed by Testo in gastrocnemius (1.34 fold amplification; Dunn’s test, p = 0.616) and in quadriceps (6.29 fold amplification; Dunn’s test, p = 0.0154).

The other component of the ubiquitin-proteasome system is the proteasome itself. In cases where E3 ligases are upregulated, it may be the case that availability of proteasomes is limiting the ubiquitin-proteasome system. However, in day 3 samples, where atrogene upregulation was at its peak, Dexa upregulated proteasome chymotrypsin-like enzymatic activity in quadriceps (18.7% increase;

a

a

a

one day

0.5

1

2

4

gastrocnemius

Trim63 mRNA

a

b

a,b

three days

0.5

1

2

4

8

16

32

gastrocnemius

Trim63 mRNA

a

a

a

seven days

0.25

0.5

1

2

4

gastrocnemius

Trim63 mRNA

a

b

a,b

0.5

1

2

4

8

16

32

64

128

256

quadriceps

Trim63 mRNA

a,b

a

b

0.5

1

2

4

8

16

32

quadriceps

Trim63 mRNA

a

a,b

b

0.25

0.5

1

2

4

quadriceps

Trim63 mRNA

a

a,b

b

0.5

1

2

4

gastrocnemius

Fbxo32 mRNA

a

a

a

0.5

1

2

4

8

16

32

gastrocnemius

Fbxo32 mRNA

a

a

a

0.25

0.5

1

2

4

gastrocnemius

Fbxo32 mRNA

Veh Dexa Comb Veh Dexa Comb

a

b

a,b

0.5

1

2

4

8

16

32

64

128

256

quadriceps

Fbxo32 mRNA

a,b

a

b

0.5

1

2

4

8

16

32

quadriceps

Fbxo32 mRNA

a

a

a

0.25

0.5

1

2

4

Veh

Dexa Comb

quadriceps

Fbxo32 mRNA

Figure 7: Changes in atrogenes MuRF-1 / Trim63 and MAFbx / Fbxo32 expression following Dexa and / or Testo treatments, in quadriceps and gastrocnemius muscles (n=3-5, normalized to GAPDH).

a

a

a

one day

0.0

0.5

1.0

1.5

quadriceps proteasome

activity (rel.u.)

a

a

a

three days

0.0

0.5

1.0

1.5

quadriceps proteasome

activity (rel.u.)

a

a

a

seven days

0.0

0.5

1.0

1.5

quadriceps proteasome

activity (rel.u.)

a

a

a

0

1

2

3

Comb

Veh

Dexa

triceps proteasome

activity (rel.u.)

a

a

a

0

1

2

3

Veh

Dexa

Comb

triceps proteasome

activity (rel.u.)

a

a

a

0

1

2

3

Veh

Dexa

Comb

triceps proteasome

activity (rel.u.)

Figure 8: Changes in proteasome chymotrypsin-like enzymatic activity following Dexa and / or Testo treatments, in quadriceps and triceps muscles (n=4-6).

Dunn’s test, p = 0.157) and triceps (111% increase; Dunn’s test, p = 0.0884; Fig. 8). This component of atrophy was also inhibited by Testo, whose co-administration reduced proteasome activity in quadriceps (24.2% reduction; Dunn’s test, p =

0.435) and triceps (30% reduction; Dunn’s test, p = 0.609) at day 3.

Overall, the good correlation with the macroscopic loss of muscle suggests that the proteasome-ubiquitin system is an effector GAML and of its alleviation by

AAS.

**Autophagy markers during dexamethasone and testosterone treatments**

Rat microarray studies indicated that a series of genes related to autophagy are upregulated by Dexa and downregulated by Testo. However, these findings have never been tested in vivo in mice. Having established a model of GAML, I investigated whether the autophagy markers are correlated with muscle loss. A panel of three genes indicated that Dexa-induced gastrocnemius atrophy is associated with a reduction in autophagy-related transcription (Fig. 9) The repression is ampler in day 3 samples, with mRNA for beclin / Becn1 reduced 9.81-fold (Dunn’s test, p = 0.0213), cathepsin L / Ctsl reduced 3.89-fold (Dunn’s test, p = 0.304), and LC3 / Map1lc3b reduced 5.61-fold (Dunn’s test, p = 0.359). Across muscle groups and time points, Testo co-administration lacked detectable

effect.

I tested the hypothesis that autophagy mediates protein degradation in GAML by measuring lysosomal enzymatic activity in muscle lysates (Fig. 10). Unexpectedly, cathepsin L enzymatic activity was reduced in all the assayed muscle, in a progressive manner. At day 1, cathepsin L activity was suppressed by Dexa in gastrocnemius (16.4% reduction; Dunn’s test, p = 0.02), quadriceps (14.5% reduction; Dunn’s test, p = 0.0294), and triceps (26.8% reduction; Dunn’s test, p = 0.00219). At day 3, cathepsin L activity was further suppressed by Dexa in gastrocnemius (19.7% reduction; Dunn’s test, p = 0.223), quadriceps (44.5% reduction; Dunn’s test, p = 0.00201), and triceps (44.5% reduction; Dunn’s test, p = 0.00107). In day 7 samples, Dexa-induced repression of cathepsin L activity reached 36.3% in gastrocnemius (Dunn’s test, p = 0.00365), 44.4% in triceps

(Dunn’s test, p = 0.0486), and 44.9% in quadriceps (Dunn’s test, p = 0.0091).

Testo co-administration had no statistically significant effect on cathepsin L

a

a

a

one day

0.125

0.25

0.5

1

2

4

8

Becn1 mRNA

a

b

a,b

three days

0.125

0.25

0.5

1

2

4

8

Becn1 mRNA

a

a,b

b

seven days

0.125

0.25

0.5

1

2

4

8

Becn1 mRNA

a

a

a

0.125

0.25

0.5

1

2

4

Ctsl mRNA

a

a

a

0.125

0.25

0.5

1

2

4

Ctsl mRNA

a

a

a

0.125

0.25

0.5

1

2

4

Ctsl mRNA

Veh Dexa Comb Veh Dexa Comb

a

a

a

0.125

0.25

0.5

1

2

4

Map1lc3b mRNA

a

a

a

0.125

0.25

0.5

1

2

4

Map1lc3b mRNA

a

a

a

0.125

0.25

0.5

1

2

4

Veh

Dexa

Comb

Map1lc3b mRNA

Figure 9: Changes in autophagy-related genes following Dexa and / or Testo treatments, in quadriceps and triceps muscles (n=3-4, normalized to GAPDH). enzymatic activity, although a trend of reversal to baseline can be seen in day 1 and 3 samples.

In literature, another line of evidence for the putative upregulation of autophagy during GAML was the increase in lipidated, fast migrating LC3 protein (also known as LC3-II). In murine muscle, detecting this form has been difficult, because it is significantly less frequent than its slower migrated counterpart (Fig. 11). In gastrocnemius at day 7, there was a trend towards enrichment of LC3-II in absolute terms upon Dexa treatment (91.6% increase, when normalized to

GAPDH; Dunn’s test, p = 1). However, LC3-II changes become negligible, when

LC3-II is normalized to its precursor, LC3-I (-0.339% increase; Dunn’s test, p = 1).

The trend towards increased absolute LC3-II in gastrocnemius is reversed by a trend towards basal levels upon Testo co-administration (29.8% decrease, when normalized to GAPDH; Dunn’s test, p = 1).

Findings like the upregulation of LC3-II may have suggested to other authors that GAML relies on autophagy. However, the overall evidence suggests that GAML is correlated with rampant autophagy downregulation, while AAS co-administration has a limited effect on autophagy. Immunoblots and enzymatic assays indicated a similar disconnection between the calpain / calpastatin system and the AAS-induced muscle sparing (Fig. 12). The ubiquitin - proteasome system emerges as the main effector of GAML and the main target of its alleviation by AAS.

## Protein synthesis modulation during Dexa-induced muscle atrophy

In order to discern possible changes in protein synthesis, I investigated a series of its translation regulators. A well-documented manner of translational shutdown upon various conditions, such as endoplasmic reticulum stress, is

a

b

a,b

one day

0.0

0.5

1.0

1.5

gastrocnemius cathepsin

activity (rel.u.)

a

a

a

three days

0.0

0.5

1.0

1.5

gastrocnemius cathepsin

activity (rel.u.)

a

b

b

seven days

0.0

0.5

1.0

1.5

gastrocnemius cathepsin

activity (rel.u.)

a

b

a

0.0

0.5

1.0

1.5

quadriceps cathepsin

activity (rel.u.)

a

b

a,b

0.0

0.5

1.0

1.5

quadriceps cathepsin

activity (rel.u.)

a

b

b

0.0

0.5

1.0

1.5

quadriceps cathepsin

activity (rel.u.)

a

b

a,b

0.0

0.5

1.0

1.5

Veh

Dexa Comb

triceps cathepsin

activity (rel.u.)

a

b

a,b

0.0

0.5

1.0

1.5

Veh

Dexa Comb

triceps cathepsin

activity (rel.u.)

a

b

b

0.0

0.5

1.0

1.5

Veh

Dexa Comb

triceps cathepsin

activity (rel.u.)

Figure 10: Changes in cathepsin L enzymatic activity following Dexa and / or Testo treatments, in quadriceps and triceps muscles (n=4-6).

D

—

—

+

+

+

+

T

—

—

—

—

+

+

LC3

GAPDH



slow >

fast >

a

a

a

0

1

2

3

LC3−II protein normalized to GAPDH

a

a

a

0

1

2

3

LC3−II protein normalized to LC3−I

Figure 11: Changes in hyperlipidated (fast-migrating) LC3 isoform following Dexa and / or Testo treatment in gastrocnemius muscles (n=4).

a

b

b

one day

one day

0.0

0.5

1.0

1.5

2.0

Dexa Comb

Veh

gastrocnemius calpain

activity (rel.u.)

a

a

a

three days

three days

0.0

0.5

1.0

1.5

2.0

Veh

Dexa Comb

gastrocnemius calpain

activity (rel.u.)

a

a

a

seven days

seven days

0.0

0.5

1.0

1.5

2.0

Veh

Dexa Comb

gastrocnemius calpain

activity (rel.u.)

Figure 12: μ-calpain enzymatic activity, during 7 days of Dexa and / or Testo treatments, in gastrocnemius muscles (n=3-6).

D

—

—

+

+

+

+

T

—

—

—

—

+

+

Ⓟ

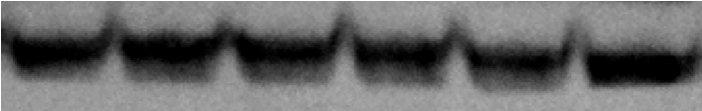
2

eIF

-

α

GAPDH



a

a

a

0.0

0.5

1.0

1.5

Veh

Dexa

Comb

phospho−eIF2alpha

(

normalized to GAPDH

)

Figure 13: Levels of phosphorylated eIF2α (Ser 51) following Dexa and / or

Testo treatment in gastrocnemius muscles (normalized to simultaneously resolved GAPDH; n=4).

phosphorylation of the initiation factor eIF2α by a diverse set of kinases. In agreement with published studies, there was no statistically significant change in the gastrocnemius of mice treated with Dexa (Dunn’s test, p = 1; Fig. 13).

Another regulator of proteins synthesis that was shown to play a role in muscle atrophy upon starvation is eIF3f. In levator muscles of mice treated with Dexa, there was no statistically significant change in the levels of eIF3f (Dunn’s test, p = 0.971; Fig. 14).

Finally, Dexa was hypothesized to achieve protein synthesis inhibition by stimulating the negative regulator 4E-BP. In Dexa-treated mice, gastrocnemius 4E-BP protein levels were upregulated (25.6% increase, when normalized to GAPDH; Dunn’s test, p = 0.428; Fig. 15, left). However, Dexa also stimulated phosphorylated, that is, inactive 4E-BP (62.5% increase, when normalized to GAPDH; Dunn’s test, p = 0.00669; Fig. 15, right).

After 7 days of Testo co-administration, levels of total 4E-BP were essentially identical to those yielded by Dexa alone (5.6% increase; Dunn’s test, p = 0.428; Fig. 15, right). At the same time, Testo co-administration was associated with

D

—

—

+

+

+

+

T

—

—

—

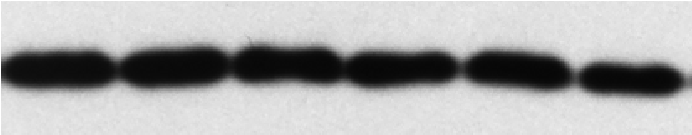
—

+

+

eIF3f

GAPDH



a

a

a

0.0

0.5

1.0

1.5

Veh

Dexa

Comb

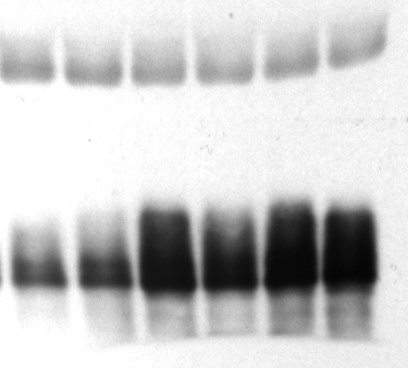
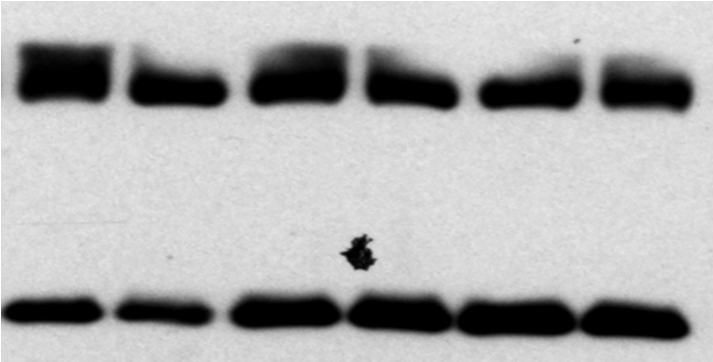
eIF3f

(

)

normalized to GAPDH

Figure 14: Levels of eIF3f following Dexa and / or Testo treatment in levator ani muscles (normalized to simultaneously resolved GAPDH; n=4).

GAPDH GAPDH

total 4EBP Ⓟ-4EBP

D — — + + + + D — — + + + +

T — — — — + + T — — — — + +

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

total4EBP

(

normalized to GAPDH

)

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

phospho 4EBP

normalized to GAPDH

)

(

Figure 15: Levels of total (left) and phosphorylated (right) 4E-BP following Dexa and / or Testo treatment in gastrocnemius muscles (normalized to GAPDH; n=4). lower phosphorylated 4E-BP (9.83% increase, when normalized to GAPDH; Dunn’s test, p = 0.49; Fig. 15, left). While it is difficult to estimate how amplitude of change in unphosphorylated, that is, active 4E-BP, the combination of increase in total 4E-BP and decrease in inactivated 4E-BP is unlikely to facilitate muscle recovery.

Overall, regulators of protein synthesis appear largely unchanged by Dexa and

Testo at later stages of their administration.

## Foxo pathway response to dexamethasone and testosterone

Previous studies on C2C12 cultured cells showed that the atrogene response is crucially regulated by Foxo transcription factors. The review section describes a series of at least five means by which Dexa is thought to stimulate Foxo transcriptional program. Most of them converge on upregulation of Foxo transcripts. I set to test the hypothesis that muscle Foxo expression is modulated by Dexa and Testo in muscle.

In this study, Foxo transcription factors were strongly induced during the early stages of GAML (Fig. 16). In quadriceps at day 1, Foxo3a was 78.1-fold upregulated (Dunn’s test, p = 0.00906). In day 3 samples, Foxo3a activation was less ample (3.17-fold; Dunn’s test, p = 0.0529). In the samples from day 7, Dexa treatment was associated with a non-significant downregulation of Foxo3a

(Dunn’s test, p = 0.175).

In contrast, Testo downregulated Foxo transcription factors at later stages of

GAML. In day 3 samples, Testo co-administration repressed Foxo3a 2.99-fold (Dunn’s test, p = 0.195). The trend was present in day 7 samples too, with Foxo3a repressed 2.04-fold (Dunn’s test, p = 0.304) by Testo co-administration. Similar patterns of modulation were seen in Foxo1 and Foxo4 transcripts (data not

a

b

a,b

one day

0.5

1

2

4

8

16

32

64

128

256

512

1024

Foxo3a mRNA

a

a

a

three days

0.5

1

2

4

8

Foxo3a mRNA

a

b

a,b

seven days

0.0625

0.125

0.25

0.5

1

2

4

Foxo3a mRNA

a

b

a,b

0.5

1

2

4

8

16

32

64

Comb

Dexa

Veh

Klf15 mRNA

a,b

a

b

0.5

1

2

4

8

Klf15 mRNA

a

a,b

b

0.0625

0.125

0.25

0.5

1

2

4

Veh

Dexa Comb

Klf15 mRNA

Figure 16: Changes in the expression of Foxo3a (top) and Klf15 (bottom) transcription factors, during 7 days of Dexa and / or Testo treatments in quadriceps muscles (n=4, normalized to GAPDH).

shown).

The changes in Foxo transcription factors were mirrored by similar changes in their positive regulator Klf15. Dexa stimulated Klf15 expression at day 1 (13.9-fold upregulation; Dunn’s test, p = 0.00906), and day 3 (2-fold upregulation; Dunn’s test, p = 0.222). Testo co-administration reversed Klf15 changes, with the strongest repression, 3.28-fold, in day 3 samples (Dunn’s test, p = 0.017).

## Akt pathway response to dexamethasone and testosterone

A alternative way to achieve Foxo modulation is through their neutralization by Akt. As described in the literature review section, activation of Akt is dependent on its phosphorylation on two residues, Ser 473 and Thr 308. I investigated the effect of the two steroids on Ser 473 phosphorylation, as this was hypothesized to be the main regulator of Foxo specificity at the time of the experiment.

After 7 days, neither drug changed the level of total Akt in levator, nor gastrocnemius (Fig. 17, left). At the same time, Dexa repressed phosphorylation of a large degree in gastrocnemius (35.6% reduction in absolute densitometry of phospho-Ser473 Akt, when normalized to GAPDH; Dunn’s test, p = 0.304), and levator ani (46.2% reduction; Dunn’s test, p = 0.146). However, Testo co-administration had no significant effect on phospho-Ser473 Akt in gastrocnemius (0.842% increase, p = 1). Interestingly, Testo co-administration repressed phospho-Ser473 in levator ani, a extremely AAS-sensitive muscle

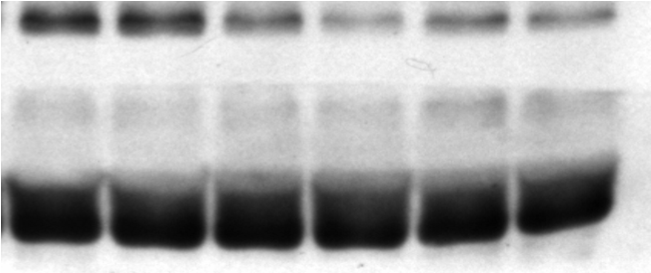
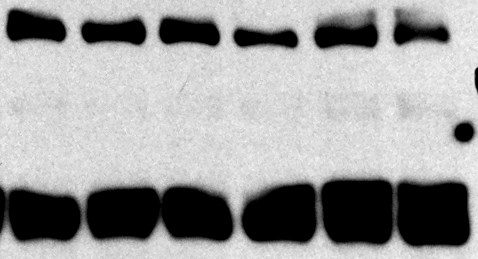
(33.1% decrease; Dunn’s test, p = 0.823).

The other relevant phosphorylation on Akt , Thr 308, is controlled by REDD1.

I investigated the effects of the two steroids on REDD1 / Ddit4 transcription (fig.

18). Dexa upregulated Redd1 expression in gastrocnemius collected at days 1

(7.48-fold; Dunn’s test, p = 0.0362), 3 (27.8-fold; Dunn’s test, p = 0.0912), and 7

 total Akt Ⓟ473-Akt

GAPDHGAPDH

D — — + + + + D — — + + + +

T — — — — + + T — — — — + +

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

gastrocnemius total Akt

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

gastrocnemius phospho−Ser473 Ak

a

a,b

b

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

gastrocnemius phospho/total Akt

D

—

—

—

+

+

+

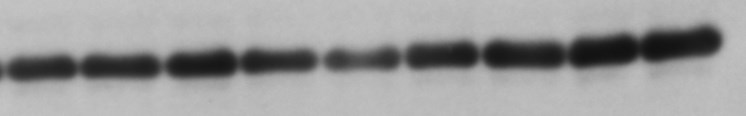
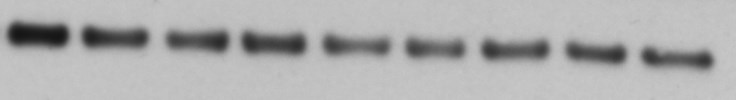
+

+

+

total Akt

GAPDH



Ⓟ

Akt

473-

GAPDH

D

—

—

—

+

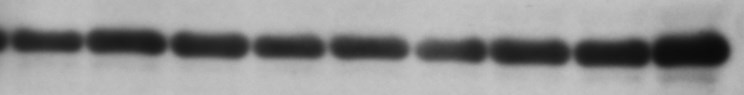
+

+

+

+

+



T — — — — — — + + + T — — — — — — + + +

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

levator total Akt

a

a,b

b

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

levator phospho−Ser473 Akt

a

a

a

0.0

0.5

1.0

1.5

2.0

Veh

Dexa

Comb

levator phospho/total Akt

Figure 17: Levels of total and phosphorylated Akt following 7 days of Dexa and / or Testo treatment in gastrocnemius (top) and levator (bottom) muscles (normalized to GAPDH; n=4).

a

b

b

one day

1

2

4

8

16

32

64

128

Dexa Comb

Veh

gastrocnemius Ddit4 mRNA

a

a,b

b

three days

1

2

4

8

16

32

64

128

Dexa Comb

Veh

gastrocnemius Ddit4 mRNA

seven days

1

2

4

8

16

32

64

128

Veh

Dexa Comb

gastrocnemius Ddit4 mRNA

Figure 18: Changes in Redd1 transcripts, during 7 days of Dexa and / or Testo treatments in gastrocnemius muscles (n=4, normalized to GAPDH).

(2.64-fold; Dunn’s test, p = 0.255). Confirming the rat microarray studies, at day

7, Testo co-administration induced a sizable repression of REDD1 (1.85-fold; Dunn’s test, p = 0.143).

## IGF-I changes during dexamethasone and testosterone administration

As explained in the review section, IGF-I was one of the few genes whose expression was changed in opposite ways by Dexa and Testo in rat microarrays. I tested the hypothesis that Dexa and Testo alter IGF-I expression in mice, in vivo (Fig. 19).

In gastrocnemius, Dexa downregulated IGF-I expression in samples collected at days 1 (4.35-fold; Dunn’s test, p = 0.0213), 3 (4.03-fold; Dunn’s test, p = 0.0105), and 7 (2.25-fold; Dunn’s test, p = 0.0362). Dexa-induced downregulation of IGF-I was also detected in quadriceps at days 3 (2.87-fold; Dunn’s test, p = 0.0319), and 7 (2.41-fold; Dunn’s test, p = 0.0213).

Testo reliably reversed this change. In gastrocnemius, Dexa downregulated

IGF-I expression in samples collected at days 1 (2.23-fold; Dunn’s test, p = 0.421),

3 (2.55-fold; Dunn’s test, p = 0.0825), and 7 (1.85-fold; Dunn’s test, p = 0.175). A

8 8 8

a

a,b

b

one day

0.125

0.25

0.5

1

2

4

quadriceps Igf1 mRNA

a

a,b

b

three days

0.125

0.25

0.5

1

2

4

quadriceps Igf1 mRNA

a

b

a,b

seven days

0.125

0.25

0.5

1

2

4

quadriceps Igf1 mRNA

a

b

a,b

0.125

0.25

0.5

1

2

4

8

Comb

Dexa

Veh

gastrocnemius Igf1 mRNA

a

b

a,b

0.125

0.25

0.5

1

2

4

8

Comb

Dexa

Veh

gastrocnemius Igf1 mRNA

a

b

a,b

0.125

0.25

0.5

1

2

4

8

Veh

Dexa

Comb

gastrocnemius Igf1 mRNA

Figure 19: Changes in the expression of IGF-I, during 7 days of Dexa and / or Testo treatments in gastrocnemius muscles (n=4, normalized to GAPDH).

similar trend was seen in quadriceps, but the amplitude was smaller, perhaps owing to the fact that Dexa-induced changed in IGF-I were smaller.

Throughout this section, the molecular correlates of GAML, as found in rats, were confirmed in this novel mouse model. However, few of the effects of Dexa were reversed by Testo. Among the putative mechanisms of alleviation of GAML, there is good evidence for the repression of the proteasome - ubiquitin system and for restoration of intramuscular IGF-I. In the next section, I describe my efforts to better understand these myoprotective mechanisms, using an in vitro model.

**6. IN VITRO FINDINGS**

**Testosterone alleviates dexamethasone-induced atrophy of cultured cells**

In mouse explanted myofibers and C2C12 cultured myotubes, muscle atrophy is a cell-autonomous phenomenon, with all its essential manifestations present in vitro. I hypothesized that the muscle protection by Testo could be replicated in

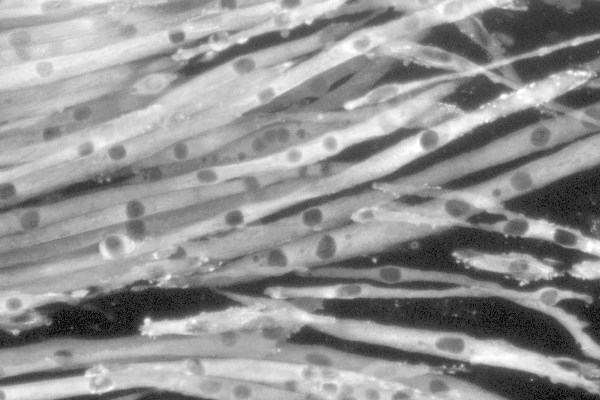
C2C12 cells. Indeed, fully differentiated C2C12 myotubes treated for 48 h with 50 µM Dexa lost 8.66% in diameter (Tukey’s HSD, p = 0.000316).

Co-administration of 300 nM Testo re-established basal diameters (Tukey’s HSD vs. Dexa alone, p = 1.02e-05).

In another experiment, I tested the ability of Testo to preserve protein content in C2C12 cells. In order to determine the time course of protein content, cells were treated with (A) vehicle, (B) 1 µM Dexa, (C) 1 µM Dexa and 100 nM Testo, or (D) 1 µM Dexa and 500 nM Testo, for up to three days, in increments of 24 h (Fig. 21, top). All the treatments led to loss of total protein starting from the third day, presumably due to senescence and / or loss of viability (ANOVA treatment x time, p = 0.000463 for time variable). When all the samples were analyzed together, treatment had no significant effect (ANOVA treatment x time, p = 0.0943 for treatment variable).

However, there were differences in total protein density prior to the third day, which mimicked the in vivo findings. These trends become apparent when data is represented after normalization to the initial time point (Fig. 21, top), or when data is analyzed only across days 1-3. In the first two days, there were no relevant changes in total protein content in cells treated with vehicle. Total protein concentration was 81.8 µg/cm2 in the first day, 81.1 µg/cm2 in the second, and

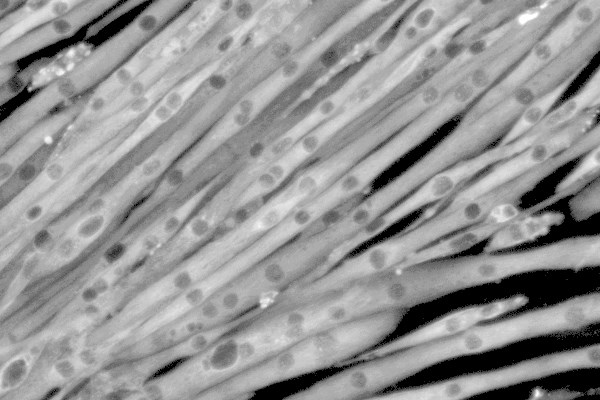
146



(

Veh

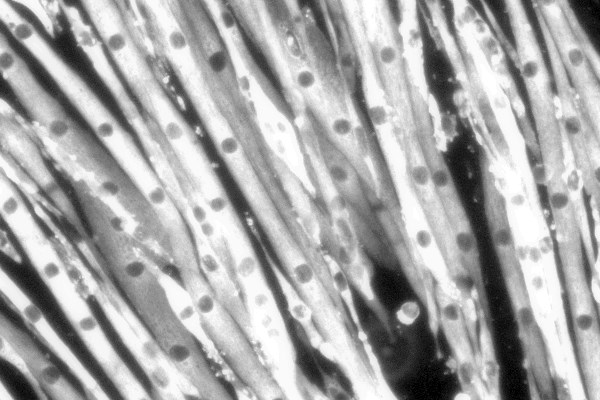
)



(

Dexa

)



a

b

a

0.0

0.3

0.6

0.9

Veh

Dexa

Comb

mean diameter (rel. u.)

(Dexa + Testo)

Figure 20: Changes in C2C12 myotube diameter following Dexa and / or Testo treatments. Micrographs are representative for cells receiving (top left) vehicle, or (top right) 50 µM Dexa, or (bottom left) 50 µM Dexa and 300 nM Testo. Bottom right image compares average myotube diameters for the three treatments (n=510-740).

65

70

75

80

85

90

2

1

3

4

day

total protein density

µg / cm

(

2)

Veh

1

µM Dexa

1

µM Dexa

+

100 nM T

1

µM Dexa

+

500 nM T

90

%

%

95

100

%

105

%

%

110

1

2

3

4

day

total protein density

normalized to initial time point

)

(

Figure 21: Changes in C2C12 myotube total protein content following Dexa and / or Testo treatments (Top) Absolute total protein density. (Bottom) The same data, after normalization to the average of the initial time point for each condition (n=5-6).

81.6 µg/cm2 in the third (ANOVA p = 0.957). In contrast, Dexa-treated cells lost protein, declining from 83.3 µg/cm2 in the first day to 81.4 µg/cm2 in the second and 78.9 µg/cm2 in the third day of the experiment (5.32% loss; ANOVA p =

0.957). Co-administration of Testo more than compensated the effect of Dexa.

Rather than losing protein, cells receiving a combination of 1 µM Dexa and

100 nM Testo gained 4.99 % total protein over the first two days of the experiment (ANOVA p = 0.957). Moreover, increasing the Testo dose to 500 nM further improved protein accretion, leading to a 7.16 % protein gain during the first two days of the experiment (ANOVA p = 0.957).

Overall, the changes in cell diameter and total protein content are consonant with protein metabolism changes observed in vivo.

**Protein synthesis in cultured cells treated with dexamethasone and**

## testosterone

In preliminary experiments, I found that tracer uptake is minimal during the first two hours. Also, tracer uptake rate became essentially equal to the rate of tracer release due to protein synthesis after the first 18 h of labeling. Therefore, I performed a series of assays where the rate of protein synthesis was estimated through the rate of tracer uptake over an intermediate 6 h (Fig. 22, top). In each case, cells were differentiated over seven days, then treated with (A) vehicle, (B)

1 µM Dexa, or (C) 1 µM Dexa and 500 nM Testo, for either 6, 24, 48, or 72 hours. The media with steroids was refreshed every 24 hours. The tracer was added with a final medium change, six hours before lysis.

After lysis, cells were assayed for protein-bound tracer. There was no effect of treatment on the amount of tracer uptook by the cells (Fig. 22, middle). To avoid the confounding influence of the atrophy, data was also analyzed as specific activity , that is, protein-bound intracellular tracer normalized to the mass of cell protein (Fig. 22, bottom). The same qualitative observations could be made after normalization. There was a significant effect of time on the rate of protein synthesis (ANOVA treatment x time, p = 2.45e-08 for time variable). The time variable was significant due to a downward trend form days 2 through 4. This trend adds to the notion that C2C12 fully differentiated myotubes quickly lose their viability. When all the samples were analyzed together, treatment had no significant effect (ANOVA treatment x time, p = 0.302 for treatment variable). There is no statistically significant difference between treatments at each time

Start pharmacological

treatments

Start differentiation

Time

)

days

(

DM

T±D

±

³

H-Phe

0

d

³

H-Phe

DM

T±D

±

1

d

DM

±

T±D

2

d

DM

±

T±D

d

3

³

H-Phe

³

H-Phe

●

●

●

●

●

●

●

●

1100

1200

1300

1400

1500

1600

1

2

3

4

day

activity in cell protein

extract (pCi/well)

●

●

Veh

1

µM Dexa

1

µM Dexa

500 nM T

+

●

●

●

●

●

●

●

●

7

8

9

10

11

1

2

3

4

day

tracer enrichment in cell

protein (fmol/g)

Figure 22: Estimates of protein synthesis rate in C2C12 myotubes treated with Dexa and / or Testo. (Top) Experimental timeline. (Middle) Amount of tracer incorporated into protein per well. (Bottom) Amount of tracer incorporated into protein per gram of total protein (n=6).

point. The amplest difference in translation rate between treatments is in day 4, when, in Dexa cells, protein synthesis rate is higher than in all other conditions (specific activity 10.6% higher than vehicle, and 13.5% higher than the combination with testosterone). The trend toward increased protein synthesis with Dexa cannot explain the observed myotube atrophy induced by Dexa. Moreover, the late onset of protein synthesis upregulation indicates that this may be a compensatory response to the earlier loss of protein.

Within the limits of the C2C12 model, protein synthesis changes do not appear to mediate Dexa-induced loss of protein, nor its alleviation by Testo.

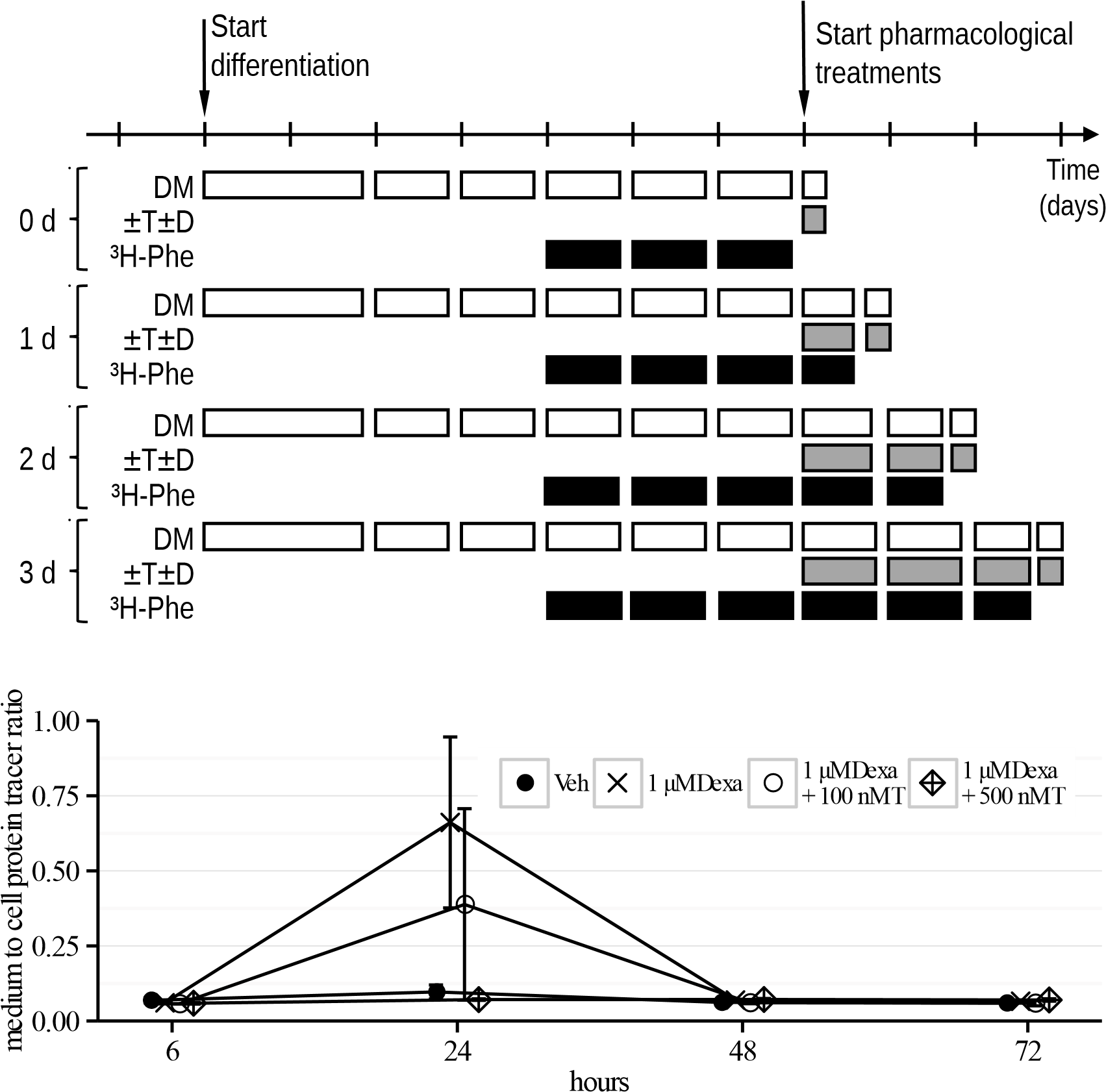
**Testosterone prevents protein catabolism upregulation induced by**

## dexamethasone

A series of complementary experiments estimated the rate of protein degradation in C2C12 myotubes. In each case, cells were differentiated over seven days, then treated with (A) vehicle, (B) 1 µM Dexa, (C) 1 µM Dexa and

100 nMTesto, or (D) 1 µM Dexa and 500 nM Testo, for either 6, 24, 48, or 72 hours (Fig. 23, top). The media with steroids was refreshed every 24 hours. The tracer was added three days before the addition of steroids, and was maintained in the media until 6 hours before lysis. Because the rate of incorporation of tracer slows down before the first 24 hours, and therefore it may be assumed that, with some approximation, the ratio of tracer to tracee in culture medium is equal to that in the pool of rapid-turnover intracellular protein.

Preliminary experiments revealed that scintillation data were not completely additive. Therefore, scintillation-based methods are semi-quantitative. The amounts of free tracer from cell extract and protein-bound tracer in the medium proteins are tens of times lower than the free tracer in the medium, and



a

a

a

a

0.00

0.25

0.50

0.75

1.00

Veh

1

µM Dexa

1

µM Dexa

+

100 nM T

1

µM Dexa

+

500 nM T

medium to cell protein tracer

ratio (24 hour)

Figure 23: Estimates of protein degradation rate in C2C12 myotubes treated with Dexa and / or Testo. (Top) Experimental timeline. (Middle) Ratio of tracer in medium to tracer in cells, across time points. (Bottom) Ratio of free tracer in medium to protein-bound tracer in cells, at day 2 (n=5-6).

protein-bound tracer in the cells. In order to avoid addition on non-additive data, protein degradation rate was primarily estimated from the ratio of free tracer in the cell culture medium to protein-bound tracer in the cell extract (Fig. 23, middle). However, the results are essentially identical when free tracer from cell extract and protein-bound tracer in the medium proteins are taken into account.

Overall, due to sample size, the treatments were not statistically significant across time points (ANOVA time X treatment, p = 0.401 for time; p = 0.259 for treatment). Nevertheless, the effects of the two steroids were ample and dose-dependent at the 24 hour time point (ANOVA between treatments, p = 0.144; fig. 23, bottom). For 24 hour vehicle-treated cells, the ratio of free medium tracer to protein-bound intracellular was 0.0968. When cells received 1 µM Dexa, the ratio of free medium tracer to protein-bound intracellular increased to 0.661 (Tukey’s HSD vs. vehicle, p = 0.202). Co-administration of Testo had a dose-dependent inhibitory effect on protein degradation. When Dexa was supplemented with 100 nM Testo, the ratio of free medium tracer to protein-bound intracellular was reduced to 0.388 (Tukey’s HSD vs. Dexa alone, p = 0.776). When Dexa was supplemented with 500 nM Testo, the ratio of free medium tracer to protein-bound intracellular was further reduced to 0.0713 (Tukey’s HSD vs. Dexa alone, p = 0.173).

A similar trend was recorded at the 48 hour time point. For 24 hour vehicle-treated cells, the ratio of free medium tracer to protein-bound intracellular was 0.062. When cells received 1 µM Dexa, the ratio of free medium tracer to protein-bound intracellular increased to 0.0699 (Tukey’s HSD vs. vehicle, p = 0.182). Co-administration of 100 nM Testo reduced the ratio of free medium tracer to protein-bound intracellular to 0.0607 (Tukey’s HSD vs. Dexa alone, p = 0.12).

Overall, Dexa myotube atrophy is correlated with an increase in tracer release, indicating an upregulation of protein degradation. Testo protection of myotubes is correlated with an inhibition of protein degradation.

**Mechanisms of androgenic myoprotection in cultured myotubes**

## treated with dexamethasone

In previous experiments, I established that loss of C2C12 myotube protein during Dexa treatment is associated with an increase in protein degradation at 24 hours after the initiation of steroid. Using a series of chemical inhibitors, I investigated the molecular mechanisms that could mediate this catabolic upregulation (Fig. 24, top). As in the previous experiment, cells were differentiated over 7 days. In the final three days of differentiation, medium with tracer was refreshed daily. For the final 24 hours, the tracer was removed, and cells were treated with (A) vehicle, (B) 100 nM Dexa, or (C) 100 nM Dexa and

300 nM Testo. In order to interfere with putative proteolytic pathways, other sets of cels were treated with (D) 100 nM Dexa and 25 µM chloroquine, an inhibitor of autophagy, or (E) 100 nM Dexa and 5 µM MG132, an inhibitor of the proteasome. Finally, in order to interfere with IGF-I signaling, another set of cells were treated with (F) 100 nM Dexa, 300 nM Testo, and 50 nM pycropodophilin, an inhibitor of IGF-1R.

The medium and the cells were fractionated as in the previous experiment. Protein degradation rate was estimated via the ratio of free tracer in the medium to protein-bound tracer in the cells. When cells were treated with vehicle alone, the ratio of free medium tracer to protein-bound intracellular was 0.826. When cells received 100 nM Dexa, the ratio of free medium tracer to protein-bound intracellular increased to 1.55 (Tukey’s HSD vs. vehicle, p = 0.000104).

Start

steroid

treatments

Start

differentiation

Time (days)

³

H-Phe

DM

±

T±D

Protein

labeling

Collection of

measured medium

a

b

a

c

a

a

0.0

0.5

1.0

1.5

Veh

100

nM Dexa

nM Dexa

100

+

25 µM CHQ

100

nM Dexa

+

5 µM MG

132

100

nM Dexa

+

300 nM T

100

nM Dexa

+

300 nM T

+

50 nM PPP

medium to cell protein tracer

ratio (24 hour)

Figure 24: Interference of protein degradation in C2C12 myotubes treated with Dexa with chemical inhibitors. (Top) Experimental timeline. (Bottom) Ratio of tracer in medium to tracer in cells (n=5-6).

The upregulation in catabolism was dependent on both proteasome and lysosome actions. When the lysosome was inhibited with chloroquine, the ratio of free medium tracer to protein-bound intracellular was brought back to basal levels (0.614 (Tukey’s HSD vs. Dexa, p = 1.44e-06). However, proteasome inhibition had a more ample result, with the ratio of free medium tracer to protein-bound intracellular depressed to 0.132 (Tukey’s HSD vs. Dexa, p = 2.4e-10).

Testo repressed the upregulation of catabolism brought by Dexa to basal levels (free medium tracer to protein-bound intracellular tracer 0.714; Tukey’s HSD vs. Dexa, p = 2.35e-05). The protective effect of Testo was not altered by the inhibition of the IGF-1R pathway (free medium tracer to protein-bound intracellular tracer 0.701; Tukey’s HSD vs. Dexa, p = 1).

Overall, in C2C12 cells, Dexa upregulates protein degradation, mainly through the activation of the proteasome. Testo reverse the activation of proteolysis in an

IGF-1R-independent manner.

**7. DISCUSSION**

**Testosterone alleviates dexamethasone-induced muscle atrophy in mice**

The present work investigated the molecular mechanisms mediating androgen alleviation of GC-induced muscle atrophy in mouse. While the myoprotective action of Testo was proven in humans and dissected in rats, it was not studied in mice. In fact, although some studies investigated transcriptional changes in mouse muscle in response to Dexa, surprisingly few describe macroscopic atrophy.

A 10 mg/(kg d) Dexa dose induced sizable loss of muscle, as evidenced by lean body mass and individual muscle weight changes. The effect was progressive, with the losses of lean body mass at day 3 being roughly half of those at day 7. The observed time dependence is evidence for a Dexa-specific effect. Moreover, lean body mass is essentially unchanged in mice treated with vehicle alone. The measurement of fat body mass indicated that the mice treated with vehicle had essentially unchanged tissue fat during the experiment. The unchanged lean and fat tissue content is evidence for the fact that the experimental manipulations had no effect by themselves. The well-being of the study animals was further proven by the fact that Dexa exerted its expected stimulating action on body fat accretion.

The change in total body mass was approximately equal to that in lean body mass. This may explain why body weight changes (Fig. 4), rather than body weight at sacrifice (Fig. 3), were significantly altered by Dexa. No other organ or tissue appeared to undergo atrophy upon Dexa treatment. Dissection revealed no significant changes in the size of viscera. No detectable changes were seen in the wet weight of the heart (data not shown).

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It is unclear what causes the variation in Dexa sensitivity between muscles. Gastrocnemius and quadriceps appeared as more sensitive muscle to Dexa, whereas tibialis was essentially unchanged. The lack of response in tibialis was surprising, given how common mouse tibialis manipulation, such as electroporation of DNA, is described in literature. However, there is no published account of GC-induced loss of tibialis mass in mouse. All three muscles contain nearly equal masses of fast- and slow-twitch fibers, at least in rat[631], thus excluding an explanation based on the type-specific differences in responsiveness. The lack of sensitivity in tibialis may be due to the fact that the tendinous component is weighing more in relative terms, compared to large fleshy muscles. There was a remarkable similarity between the in vivo and in vitro findings. Whereas quadriceps and gastrocnemius lost 2-3.5% of their weight for every day of Dexa, C2C12 myotubes lost about 5% of their total protein upon Dexa treatment. This rate is similar to that indicated by Desler for C2C12 myotubes differentiated over three days[559].

In contrast to the in vivo studies, cells ability to thrive degraded towards their third day of Dexa treatment, that is, their ninth day of differentiation. The overarching manifestations of cell senescence prevented the assessment of myotube atrophy beyond the 48-hour time point. Moreover, the atrophic fibers’ diameter becomes comparable with that of the nucleus (Fig. 20). Further reductions in cell diameter would have required a shrinkage of the nucleus, which has not been observed in the first two days of myotube atrophy. It was therefore impossible to develop a more chronic in vitro model of muscle atrophy.

Co-administration of Testo alleviated all macroscopic Dexa effects. Similar to the profile of Dexa action, in absolute terms, the recovery in lean body mass was approximately equal to that in total body weight. The percentage by which total body weight, lean body mass, and individual muscles recovered are similar, indicating that Testo action was also limited to muscle.

The experimental protocol was not ideal for observing the time course of the

Testo action. On one hand, the body weight changes during the 7 day experiment (Fig. 4, top) suggests that the protective action of Testo starts early on, from the first day of experiment. On the other, the time course is not confirmed by the evolution of individual muscles’ weight (Fig. 6), where 7-day samples appear effectively protected by Testo, whereas 3-day samples show a more limited anabolic response. This discrepancy originates from the fact that the mice analyzed in the 7-day study were slightly more developed than those used in the 3-day sample, as indicated by the difference in levator muscles at sacrifice between vehicle-treated animals at each time point. Moreover, the detection of the Testo myoprotective effect may be strained by its incomplete nature. Because Dexa effect is progressive, its amplitude in the early stages is necessarily small. When Dexa-induced atrophy is hard to detect, its incomplete reversal will be even harder to demonstrate. One cannot exclude a temporal dissociation between the actions of the two steroids, but the changes surrounding the acute onset of GAML were not investigated in more detail, as they were beyond the scope of this work. At the 7-day time point, which is the most accurate model of chronic steroid myopathy, the alleviation of GAML is well-established.

The alleviation following Testo was incomplete in terms of lean body mass and individual muscle weights. In contrast, total body weight completely recovered. The source of this discrepancy is not known. Dissection revealed no other viscera with an appearance of hypertrophy following Testo co-administration. The dose of Testo used here was shown to be effective in rats.

However, the Dexa dose used in the rat studies was much smaller, suggesting that mice studied here would have benefited even more from an increased dose of Testo. Moreover, mice may be intrinsically less responsive to Testo. The effective dose of Testo for fully preventing GAML should be the aim of future studies. Within the limits of present data, it may be the case that no Testo dose would overcome Dex-induced atrophy, which would then imply that Dexa pro-atrophic mechanisms include pathways outside the scope of AAS action.

There were differences in the responsiveness to Testo between muscle groups. Similar to Dexa sensitivity, an important component appears to be the tendinous content. Levator ani is more responsive to Testo than to Dexa, owing probably to the increased presence of AR compared to other muscles[632]. The apparent dependence of Testo response amplitude to the expression of AR indicates that alleviation of GAML by AAS is a specific effect, rather than an interaction at GR

level.

In vitro, myoprotective action of Testo was similarly present at late time-points. The protection of C2C12 myotubes was dose-dependent, based on the total protein assay. In a parallel to the in vivo system, the myoprotective effect was more salient when data was normalized to the first time point in the experiment (fig. 21 top versus bottom).

## Testosterone’s myoprotective action is based on the inhibition of the dexamethasone-induced proteasome upregulation

The mouse model of GAML largely replicated what was known from rat experiments, where a strong upregulation of the proteasome system is seconded by alterations in protein synthesis.

In vivo, the E3 ligases also known as atrogenes were upregulated. The upregulation of MuRF-1 was precocious, strong, and reached statistical significance despite the small sample size. Upregulation of MuRF-1 was confirmed at early stages in gastrocnemius, quadriceps, and triceps (Fig. 7; the latter not shown). In agreement with published studies, the more specific ligase, MAFbx, was upregulated by a lesser percentage than MuRF-1. In combination with the practical limitations on sample sizes, the low-grade activation of MAFbx strained its detection to a statistically significant degree in this experiment. Nevertheless, the invariable trend across muscle groups towards increased MAFbx expression in response to Dexa indicates that the second atrogene was also part of the atrophic transcriptional program. The massive proteasome upregulation observed in day 3 samples increased the probability of the hypothetical scenario in which protein degradation is limited by proteasome availability, rather than E3 ligases. However, day 3 also marks the peak of proteasome catalytic activity (Fig. 8). The synchronized stimulation of the proteasome and of the atrogenes indicates that the proteasome - ubiquitin system is at the center of GAML.

All these catabolic catabolic changes were inhibited by Testo co-administration. In vitro, atrogenes expression was reduced to basal levels, and, in the case of day 7 samples, even further below. The inhibitory effect of Testo on MuRF-1 was predicted by the microarray study on rat muscle. Similar to the rat study, the changes in MAFbx were also in agreement with the phenotype, but of a lesser amplitude. In addition, the proteasome activity was suppressed at its day 3 peak. This aspect of muscle atrophy, already demonstrated as a component of male post-castration muscle atrophy[622], has never been investigated in GAML, nor in its alleviation by Testo.

Upregulation of proteasome activity, independent of atrogene status, has been reported in others[438]. It is unclear how Dexa achieves this upregulation. In this experiment, limited exploration found no upregulation of transcripts for proteasome subunits A6, B10, or D4 (data not shown).

In vitro, proteasome inhibition led to nearly complete suppression of proteolysis, beyond the basal levels seen in vehicle-treated cells. This finding indicates that most of the proteolytic activity in C2C12 cells is dependent on the proteasome.

A study by Baehr and colleagues[432], which was published while my experiments were under way, showed that MAFbx knockout did not reduce the amplitude of GAML, while MuRF-1 knockout reduced GAML to about half of its amplitude in wild-type mice. My data show that, instead of rejecting the proteasome-centered model of GAML, future studies should focus on finding alternative ways by which the proteasome performs its catabolic function. For example, the simplest scenario fitting today’s data is one where the proteasome performs the clearance of the bulk of dispensable proteins. An alternative explanation, which takes into account the subsiding evolution of the atrogene surge, is that atrogenes ubiquitinate, and target for degradation, an yet unidentified myoprotective intracellular factor, thus unleashing a cascade of proteasome-independent mechanisms. Some limited attempts to detect multiple ubiquitination states for MuRF-1’s putative substrate, myosin heavy chain, have failed (data not shown). If MuRF-1 has only this limited and limiting action, disruption of MuRF-1 in Baehr’s knockout mice could have been supplanted by partially homologous genes, such as MuRF-2 or Fbxo40[633, 442]. The distinction between the two scenarios is difficult, especially in the in vivo approach. In either scenario, the role of the proteasome is indispensable for GAML. Proteasome inhibition by Testo co-administration emerges as an important mechanism of GAML alleviation.

The experiments performed in this study clearly exclude a role for autophagosome - lysosome system in digestion of bulk myofiber proteins. The in vivo data indicate a persistent suppression of autophagy-related genes across muscle groups and time points. Moreover, lysosome-associated cathepsin L enzymatic activity is suppressed by Dexa in a statistically significant manner at all time points. The downregulation of cathepsin activity and expression was more ample than the the loss of muscle protein, indicating that an active process of cathepsin degradation is activated by Dexa in vivo.

The finding of Dexa-induced autophagy downregulation was unexpected.

Some atrophic conditions, most notably starvation, lead to autophagy regulation. The rat microarray findings found cathepsin L among the set of genes upregulated by Dexa. Even in this study, a narrow measurement, the accumulation of lipidated LC3 appeared upregulated by Dexa (Fig. 11, middle). However, the microarray results were never validated by qRT-PCR. In the present work, when the hyperlipidated LC3 form is normalized to its precursor, its levels become essentially unchanged in response to Dexa (Fig. 11, right). The accumulation of LC3 protein, both in precursor and mature form, indicates reduced capacity in the autophagolysosome compartment, especially in the present context of downregulated LC3 protein expression. This line of evidence corroborates with the downregulated enzymatic activity to refute a putative role for autophagy in bulk GAML.

Intuitively, it is more likely for bulk protein catabolism to be mediated by the smaller proteasome and atrogenes than the bulky autophagosome. However, in the cell culture experiments, inhibition of lysosomes had a significant protective effect, but not as ample as that of proteasome inhibition. Even in the in vivo studies, the amplitude and invariability of autophagy inhibition suggests it may have a regulatory role. While the present data solidly exclude a role for bulk protein digestion, further studies are needed to elucidate which proteins are spared from autophagy during GAML, and what is the regulatory effect of their sparing.

Prior to this work, the calpain system was the least likely effector of GAML. In agreement with the literature, this study could not substantiate Dexa-induced changes in calpain enzymatic activity, calpain, or calpastatin protein levels (data not shown). Overall, the absence of Dexa-induced amplification in catabolic activity in the cathepsin and calpain pathways reduces the scope for AAS myoprotection through inhibition of these pathways. Moreover, Testo had no reliable effect on the Dexa-induced changes in the autophagosome - lysosome pathway. Therefore, Testo protection is not mediated by inhibition of cathepsin or calpain.

Studies on protein synthesis rate have been strained by the limited technical abilities of measuring protein synthesis in mice. This study did not measure protein synthesis directly in vivo. The measurements of protein synthesis rate in vitro failed to identify any significant change in response to Dexa. C2C12 cells are surprisingly dependent on protein synthesis, with either translational inhibitor cycloheximide and puromycin leading to cell death and detachment in hours (data not shown). Overall the lack of detectable changes in protein synthesis rate agrees with findings in rat L6 cells[406] and explanted muscle[329].

In order to detect subtle changes in protein synthesis, I investigated a series of its regulators. I failed to identify changes in levels of phosphorylated eIF2α and eIF3f in response to either Dexa or Testo (Figs. 13, 14). I could not detect ATF4 protein in muscle lysates (data not shown). In the 7-day samples, both total and phosphorylated 4E-BP were upregulated by Dexa. With these data, it was unclear whether the active negative regulator of protein synthesis, unphosphorylated 4E-BP was increased or decreased by chronic Dexa. The changes in total and phosphorylated 4E-BP induced by Testo are small, and cannot lead to increased inactivation of 4E-BP. Therefore, there was no evidence that, at 7-day time point, the protective action of Testo benefits from increased protein synthesis.

Moreover, while the reported experiments were ongoing, Baehr and colleagues reported that Dexa decreased protein synthesis rate at day 3 and increased it at day 14 in mouse triceps[432]. Baehr et al. findings suggest that acute Dexa represses protein synthesis, whereas chronic Dexa is associated with a compensatory restoration of translational capacities. At the 7-day time point analyzed in this study, a measurement of protein synthesis by Baehr’s method would have been indecisive, as the muscle would have been midway in the switch from low translation to high translation rate. It is unclear what co-administration of Testo would have induced in adapting translation machinery.

In conclusion, Testo induced muscle protection through multiple mechanisms, among which inhibition of the proteasome system stood out by amplitude and persistence.

**IGF-1R signaling during dexamethasone and testosterone administration**

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## Final considerations

This work demonstrates that Testo protects mouse muscle in vivo and in vitro by reversing the Dexa-induced upregulation of proteasome. In contrast to the expectations set by the microarray studies, there was no evidence for bulk protein digestion through the autophagosome. However, Dexa and Testo significantly modulate autophagy, possibly achieving regulatory effects. The role of autophagy in GAML and its reversal by AAS must be clarified by studies on LC3-GFP mice. Dexa and Testo modulate in opposite manner the transcription of Foxo and Klf15 transcription factors. Moreover, the two steroid hormones regulate the mTOR inhibitor REDD1. More transgenic studies are required to establish the relative importance, and the eventual interaction between Foxo and mTOR pathways.

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**8. CURRICULUM VITAE**

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## Education

* PhD candidate, Boston University School of Medicine, expected graduation

2015

* BS (Licentiat in Biofizica), Universitatea Bucuresti, 2005
* BM\DM (Doctor in Medicina), Universitatea de Medicina si Farmacie Carol

Davila Bucuresti, 2002

## Research experience

* 2012-2015: Department of Medicine, Boston University School of Medicine,

P.I. Dr. Shalender Bhasin, under the supervision of Dr. Carlo Serra

**–** Subject: Testosterone alleviating glucocorticoid-induced muscle loss.

* 2009-2012: Department of Biophysics, Boston University School of

Medicine, P.I. Dr. Assen Marintchev

**–** Subject: Interactions between translation initiation factors eIF1A and eIF5B

* 2005-2008: Center for the Study of Brain, Mind and Behavior, Princeton University, P.I. Dr. Anne Treisman, FRS

**–** Subject: Brain mechanisms for statistical processing of visual scenes

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* Summer 2004: Biology Department, Universitatea Bucuresti, P.I. Dr.

Gordon Reid, under the supervision of Dr. Iurie Barbu

**–** Subject: The mediation of thermoception by ionic membrane channels

* 2001-2002: Biophysics Department, Universitatea de Medicina si Farmacie

Carol Davila Bucuresti, P.I. Dr. Dan Eremia, under the supervision of Dr.

Eva Katona,

**–** Subject: The effect of non-ionizing radiation of the mobility of the cell membrane lipids

## Peer-reviewed publications

1. Serra C, Tangherlini F, Rudy S, Lee D, Toraldo G, Sandor NL, Zhang A, Jasuja R, Bhasin S. *Testosterone improves the regeneration of old and young mouse skeletal muscle*. J Gerontol A Biol Sci Med Sci. 2013 Jan;68(1).
2. Serra C, Sandor NL, Jang H, Lee D, Toraldo G, Guarneri T, Wong S, Zhang A, Guo W, Jasuja R, Bhasin S. *The effects of testosterone deprivation and supplementation on proteasomal and autophagy activity in the skeletal muscle of the male mouse: differential effects on high-androgen responder and low-androgen responder muscle groups*. Endocrinology. 2013 Dec;154(12).
3. Guo W, Bachman E, Vogel J, Li M, Peng L, Pencina K, Serra C, Sandor NL, Jasuja R, Montano M, Basaria S, Gassmann M, Bhasin S. *The effects of short-term and long-term testosterone supplementation on blood viscosity and erythrocyte deformability in healthy adult mice*. Endocrinology. 2015

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268 **Other scientific communications**

1. Sandor NL, Hendrickson E, Sandor D, Wagner G, Pestova TV, Marintchev A. *Interplay between intra- and intermolecular interactions involving human eIF1A and eIF5B*. Abstract presented at the 2010 Meeting of Translational Control, Sept. 2010, Cold Spring Harbor, NY.
2. Sandor NL, Lee D, Toraldo G, Zhang A, Jasuja R, Bhasin S, Serra C. *The role of testosterone on the control of muscle protein synthesis and degradation*.

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1. Serra C, Lee D, Sandor NL, Toraldo G, Jang H, Jasuja R, Bhasin S.

*Characterization of the neuromuscular junction in castrated male mice*.

Poster presented at ENDO2013, The Endocrine Society’s 95th Annual Meeting & Expo, 2013, San Francisco, CA.

1. Sandor NL, Jasuja R, Serra C, Bhasin S. *Testosterone alleviates glucocorticoid myopathy by inhibiting the proteolytic machinery*. Poster presented at the 2013 Evans Center Days, Nov. 2013, Boston, MA.