

Activating Brown Adipose Tissue through Exercise (ACTIBATE) in Young Adults:

Rationale, Design and Methodology

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LIST OF ABBREVIATIONS

ACTIBATE: Activating Brown Adipose Tissue through Exercise

BAIBA: β -aminoisobutyric acid

BAT: Brown adipose tissue

BMI: Body mass index

BRITE: Brown-in-white

CAMD: *Centro Andaluz de Medicina del Deporte*

CIT: Cold-induced thermogenesis

DXA: Dual Energy X-ray Absorptiometry

¹⁸F-FDG: ¹⁸F-fluorodeoxyglucose (¹⁸F-FDG)

HRres: Heart rate reserve

iMUDS: *Instituto Mixto Deporte y Salud*

MIT: Meal-induced thermogenesis

PET/CT: Positron emission tomography/computed tomography

RCT: Randomized controlled trial

REE: Resting energy expenditure

RM: Repetition maximum

SNS: Sympathetic nervous system

UCP-1: Uncoupling protein 1

VAS: Visual analogue scales

ABSTRACT

Aims: The energy expenditure capacity of brown adipose tissue (BAT) makes it an attractive target as a therapy against obesity and type 2 diabetes. BAT activators namely catecholamines, natriuretic peptides and certain myokines, are secreted in response to exercise. ACTIBATE will determine the effect of exercise on BAT activity and mass measured by positron emission tomography/computed tomography (PET/CT, primary outcome) in young adults. ACTIBATE will also investigate the physiological consequences of activating BAT (secondary outcomes).

Methods: ACTIBATE will recruit 150 sedentary, healthy, young adults (50% women) aged 18-25 years. Eligible participants will be randomly assigned to a non-exercise group (n=50) or one of two exercise groups (n=50 each). Participants in the exercise groups will perform aerobic and strength training 3-4 days/week at a heart rate equivalent to 60% of heart rate reserve (HRres), and at 50% of 1 repetition maximum (RM) for the moderate-intensity group, and at 80% of HRres and 70%RM for the vigorous-intensity group. Laboratory measures completed at baseline and 6 months include BAT activity and mass, resting energy expenditure, meal and cold-induced thermogenesis, body temperature regulation and shivering threshold, body composition and cardiovascular disease risk factors. We will also obtain biopsies from abdominal subcutaneous white adipose tissue and skeletal muscle to analyse the expression of genes encoding proteins involved in the thermogenic machinery.

Discussion: Findings from ACTIBATE will have significant implications for our understanding of exercise and its protective effects against the development of type 2 diabetes, obesity and related metabolic diseases.

ClinicalTrials.gov ID: NCT02365129

Key words: Brown adipose tissue; energy metabolism; thermogenesis; body temperature regulation; obesity; type 2 diabetes.

INTRODUCTION

Both obesity and diabetes are serious health problems. Both unhealthy diets and physical inactivity are the main causes of the increasing levels of obesity and insulin resistance, which in turn lead to compensatory hyperinsulinaemia and ultimately type 2 diabetes. Although much research has been done, it is evident that our current knowledge is insufficient and new approaches have to be harnessed and exploited.

The ability to oxidize glucose and lipids and to dissipate energy as heat makes brown adipose tissue (BAT) an attractive target for anti-obesity and type 2 diabetes [1]. BAT is highly regulated by the sympathetic nervous system (SNS) to increase body temperature when mammals are exposed to temperatures below thermoneutrality. The heat production is mediated by uncoupling protein 1 (UCP-1), an inner-membrane mitochondrial protein exclusively expressed in BAT [1].

In 2009, a number of human studies showed that BAT exists and that is thermogenically active in adults [2-5]. More recently, brown adipocyte-like cells in white adipose tissue of both rodents and humans have been found. These cells, so called brown-in-white (BRITE, or beige) adipocytes, possess a multilocular morphology and express the brown adipocyte-specific UCP-1 [6, 7].

A potential clinical implication of activating BAT relates to the stimulation of resting energy expenditure (REE), meal-induced thermogenesis (MIT) [8] and cold-induced thermogenesis (CIT) [9]. In humans, MIT seems to be higher in those possessing BAT [10]. CIT has been strongly correlated with BAT, and it has been estimated that BAT increases total energy expenditure on 200-400 kcal/day when humans are under cold conditions [11]. In addition, it has been proposed that even if BAT activation would not produce enough energy expenditure to induce a negative energy balance, it could exert metabolic benefits on the prevention and treatment of type 2 diabetes and associated dyslipidaemia [12-14].

The SNS, through adrenergic signalling, is the classical regulator of BAT; however, recent findings have shown a pool of novel BAT activators that sidestep the need for stimulating the SNS, such as cardiac natriuretic peptides [15], irisin [16], β -aminoisobutyric acid (BAIBA) [17], interleukin-6 (IL-6) [18], and fibroblast growth factor 21 [19]. Of interest is that both SNS and non-SNS BAT activators are sensitive to exercise [20, 21]. Several studies conducted in animal models have reported an increased BAT activity and browning after an exercise-based intervention [16, 22-25], yet there is still some controversy [26-30]. In humans, there seem to exist a positive association of exercise with BAT and browning activity [31-33].

The primary objective of the ACTIBATE randomized controlled trial (RCT) is to quantify the dose-effect of different exercise intensities (i.e. no exercise, moderate-intensity and vigorous-intensity), on BAT activity and mass (primary outcome), and on REE, MIT, CIT, body temperature regulation and shivering threshold, body composition and cardiovascular disease risk factors in young adults. We will also obtain biopsies from abdominal subcutaneous white adipose tissue and skeletal muscle to analyse the expression of genes encoding proteins involved in the thermogenic machinery.

METHODS

The present study is a RCT (ClinicalTrials.gov ID: NCT02365129). The Human Research Ethics Committee of both University of Granada (n° 924) and *Servicio Andaluz de Salud (Centro de Granada, CEI-Granada)* approved the study design, study protocols and informed consent procedure. All participants will have to provide a written informed consent. Participants will be randomly allocated to the usual care (control), moderate-intensity exercise or vigorous-intensity exercise groups, and will be followed for 6 months during the exercise interventions. All the baseline and follow-up examinations will be performed in the same setting [*Instituto Mixto Deporte y Salud (iMUDS)* at the University of Granada and *Hospital Universitario Virgen de las Nieves, Granada, Spain*] and by the same investigators. The study will be performed following the ethical guidelines of the Declaration of Helsinki, last modified in 2013.

Participants and selection criteria

Participants will be students from the University of Granada (Granada, Spain). The University of Granada has $\approx 65,000$ graduate students, of which $\approx 40,000$ are 18-25 years. Eligible participants should be 18-25 years old and should have a body mass index (BMI) between 18.5 to 35 kg/m². The inclusion and exclusion criteria are listed in Table 1. We decided to implement the intervention on young adults with a wide range of BMI because overweight and obesity at this age is associated with an increased risk of cardiovascular disease among future middle-aged adults [34]. Moreover, BAT seems to be related to age and BMI [4, 35], being higher in younger and thinner people. Thus, people aged 18-25 years old are expected to present detectable BAT activity and mass [5, 36]. Further, persons of this age are legally able to sign themselves informed consent to participate in ACTIBATE, and no ethical issues are raised due to participant's age and type of measurements to be undertaken in the study. Risks will be minimized by ruling out contraindications to the testing and training protocols via a health

history and a thorough physical examination prior to the testing sessions. If any medical problems arise during the project, participants will be referred for medical evaluation and, if necessary, dropped from the study. ACTIBATE has subscribed private liability insurance covering participants, investigator's responsibility as well as the responsibility of any person involved in the study, provided that there is proper adherence to the protocol.

Table 1. Selection criteria.

Inclusion criteria	Exclusion criteria
<ul style="list-style-type: none"> -Age: 18-25 years. -BMI: 18.5-35 kg/m². -Not engaged in regular physical activity >20 min on >3 days/week. -Not participating in a weight loss program. -Stable weight over the last 3 months (body weight changes <3 kg). -Normal electrocardiogram. -Participants must be capable and willing to provide consent, understand exclusion criteria and accept the randomized group assignment. 	<ul style="list-style-type: none"> - History of cardiovascular disease. - Diabetes or hypertension. - Pregnancy, or planning to get pregnant during the study period. -Medication for hypertension, hyperlipidemia, hyperuricemia or other illness. -Beta blockers or benzodiazepines use. -Smoking. -Frequent exposure to cold temperatures (Granada is surrounded by high mountains where people can ski or do trekking). -Taking medication for thyroid. -Other significant medical conditions that are life-threatening or that can interfere with or be aggravated by exercise. - Unwillingness to either complete the study requirements or to be randomized into control or training group. - A first-degree relative with history of cancer.

Information period: participants will visit the iMUDS in two occasions. In the first visit, participants will receive a thorough explanation about the study aims, measurements to be undertaken, study requirements of the participants, and intervention types. What participants can expect from the study will be clarified, and any questions will be answered. Potential interested participants will therefore receive a short web-based questionnaire to obtain information about age, weight, height, physical activity, weight during the last 3 months, current medical history and medication use, smoking and alcohol habits, and residence.

Individuals who remain eligible after the first screening and express willingness to participate in the study will be invited to a second orientation session. In the second visit, participants will receive detailed written information about the study. Eligible participants who wish to participate in further screening will be asked to sign the informed consent. Participants will then be scheduled for their first baseline measurement visit. Figure 1 illustrates the participant flow from recruitment to randomisation.

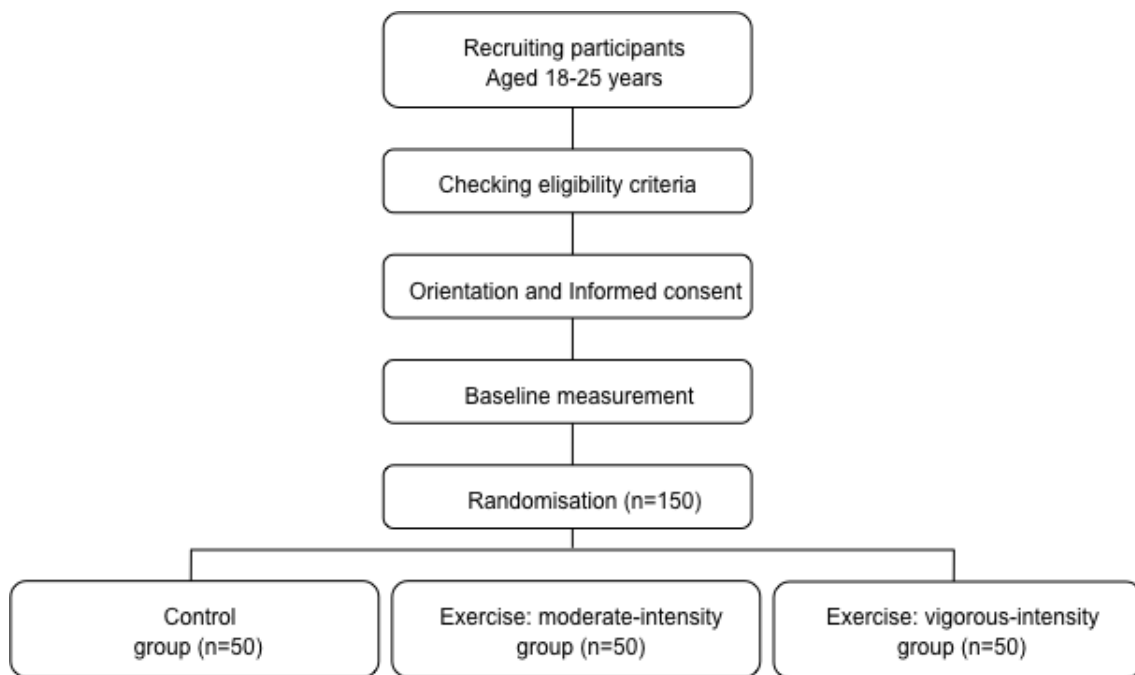


Figure 1. Flow diagram of the study participants.

Randomization and blinding

Eligible participants will be randomly assigned after completing the baseline measurements to either the control or exercise training groups. The randomization will be computer generated. Assessment staff will be blinded to participant randomisation assignment. Participants will be explicitly informed on the group to which they will be assigned as well as on the study hypotheses and will be reminded frequently not to disclose their randomisation assignments to assessment staff in the follow-up measurements. For practical and feasibility reasons, the study will be conducted in waves of maximum 15 persons.

Sample size

The estimations of the sample size and power of our study are made based on diverse observations from acute-cold exposure studies in humans, and taking assumptions extrapolated from animal models [3, 8, 37]. We used a conservative approach to sample size estimation and assumed a relatively large standard deviation based on the heterogeneity of data published in humans. Moreover, due to the nature and duration of the study as well as the complexity of the outcomes to be measured, there were conservative estimates of effect sizes, accounting for higher drop-outs than that achieved in prior exercise-based RCTs. Power calculations are made for the primary outcome measures of BAT activity and mass.

It has been estimated that 50 g of activated BAT translates to $\approx 5\%$ of REE in humans [3, 37]. An example of an ACTIBATE participant would be a sedentary man aged 19 years, body weight 90 kg, height 1.75m, and a body mass index of 29.4 kg/m^2 , with a REE of ≈ 2000 kcal/day. We anticipated increases of 10% and 20% on activated BAT mass at 6 months in moderate-intensity and vigorous-intensity exercise groups, respectively, from a baseline level of 50 g, and change score SD of 50 g. This increase could translate into a REE of 110-120 kcal/day. Affecting energy balance by 50-100 kcal/day could prevent weight gain in the vast majority of the population [38]. Assuming a bilateral alternative (which means that an effect in

either direction will be interpreted), we can detect differences of at least 10% in BAT mass with a power of >80% and α of 0.05 in a group of 17 participants. With this number of participants per group we can also detect differences of 10-20% in BAT activity with a power of >80% and α of 0.05. Since several studies reported sex-differences in BAT mass [4], all the analysis will be conducted in men and women separately. Therefore, to avoid loss of statistical power, a total of 34 (50% women) participants should be enrolled in each group. Assuming a maximum loss at follow-up of 30%, we decided to recruit 50 participants (50% women) for each of the study groups: control, moderate-intensity and vigorous-intensity groups. A total of 150 participants (of which 75 will be women and 75 men) will be enrolled in ACTIBATE. We believe this sample size is feasible and realistic based on our previous experiences in RCTs [39, 40]. We used IBM-SPSS Sample power software (version 3.0.1) for calculations.

Statistical analysis

All outcome variables will be checked for normality and results will be expressed as mean and SD or median and ranges. For between group comparisons at baseline (control group *vs* moderate-intensity and vigorous-intensity), we will analyse continuous variables with one-way analysis of variance or the non-parametric method of Kruskal-Wallis, and Chi-square tests (or exact techniques if needed), as appropriate. We will use General Lineal Models to assess the training effects (time \times group interactions) on the primary and secondary study outcomes. We will adjust multiple comparisons for mass significance [41]. We will also examine the differences between drop-outs and participants who remain in the study. We will analyse the data according to the intention-to-treat principle [42], and will handle missing data due to drop-outs or non-compliance using multiple imputation methods. To fully appreciate the potential influence of missing responses, we will perform sensitivity analysis. In addition, multiple regression analysis will be performed to evaluate potential independent predictors of BAT activity.

Participant retention and adherence

Participants will be allowed to withdraw at any time; however, to reduce participants drop out and to maintain adherence to the training program, several strategies will be used (see below). In anticipation of private commitments, vacations, etc. that might interfere with a participant's availability to come to the exercise centre for an exercise session, participants will be allowed to do exercise out of the centre. We will allow participants to do a maximum of 15 unsupervised exercise sessions throughout the study. In this situation, participants will receive a heart rate monitor and instructions on how to operate it, a kit of elastic bands to perform the strength training and the session planning. They will be asked to complete a detailed log sheet to record the type of activity completed, duration of exercise, and distance travelled during exercise, if applicable.

All sessions will be accompanied with music that participants will choose, and will be performed on an airy, well-lighted exercise room. Qualified fitness specialists will carefully supervise every training session and will work with groups of no more than 10 persons to ensure that participants are performing the exercises correctly, and at proper intensity. The training specialist and other study staff will constantly support participants.

Exercise program rationale and selection of doses

With the final aim of making the exercise program transferable to society, the basis for the specific exercise dose in ACTIBATE is the physical activity recommendations for adults proposed by the World Health Organization [43]. Since there is no information regarding the ideal exercise model to activate and recruit BAT, ACTIBATE will combine both aerobic and resistance training. A major objective of the study is to evaluate various exercise intensity levels (moderate and vigorous) that fall within the current public health recommendations to test whether higher intensity levels provide more benefit than the standard moderate-intensity level.

The length of the trial is 6 months based on results from previous large scale RCT, and based on the fact that substantial physiological adaptations occur within the first 3-6 months of exercise [44]. We have also considered the increased logistical and participation burdens, which indeed might lead to poorer adherence, as well as the cost of running a highly controlled-laboratory based study for a longer period. No dietary prescription or instructions will be provided to the participants in both control and exercise groups, except for the prescription to not change their dietary habits during the study period.

Volume. An important goal in the development of exercise doses of ACTIBATE was to keep the total volume (in minutes) of weekly exercise similar in both intervention groups while ensuring that the aerobic, as well as the resistance training, prescriptions met current guidelines [43, 45]. We will achieve both goals since the total time of aerobic exercise in both moderate-intensity and vigorous-intensity groups will be 150 minutes/week, whereas the time needed to complete the resistance training exercises will be ≈ 80 minutes for both groups.

It is estimated that 150 minutes/week of moderate-intensity [≈ 3 -5.9 metabolic equivalents (METs), 1 MET= 3.5 ml O₂/kg/min] aerobic physical activity is equivalent to 1000 kcal/week, which is associated with lower rates of cardiovascular disease and premature mortality [46]. An energy expenditure of 1000 kcal/week can also be achieved with ≈ 75 minutes/week of vigorous intensity (≥ 6 METs).

We had discussions on whether the total volume of energy expended in both moderate-intensity and vigorous-intensity groups should be equal, so that it would be possible to test the independent contributions of volume expended versus intensity. The volume selected for ACTIBATE is based on the minimum weekly physical activity time recommended by the public health organizations, with the final goal of making the results of the intervention easily transferable and understandable to the population in terms of time in minutes/week, intensity, and frequency. The lack of time is one of the main reasons reported to justify not to engage in

physical activity. For this reason, we decided to test two different training strategies that take the same time, but that due to the different intensities might lead to different results. Thus, both the moderate-intensity and vigorous-intensity groups will perform 150 minutes/week of aerobic exercise and ≈ 80 minutes/week of strength training. For the aerobic exercise, the vigorous intensity group will perform 75 minutes/week at moderate intensity (i.e. 60% of heart rate reserve (HRres)) and 75 minutes/week at vigorous intensity (i.e. 80% HRres), while the moderate-intensity group will perform the total of 150 minutes/week of aerobic training at 60% HRres. The strength training will be performed at 50% of 1 repetition maximum (RM) for the moderate-intensity group and at 70% RM for the vigorous-intensity group.

Therefore, the energy expenditure associated to the aerobic training prescribed for the vigorous-intensity group will be ≈ 1500 kcal/week (75 minutes/week of vigorous-intensity = 1000 kcal/week + 75 minutes/week of moderate-intensity = 500 kcal/week). We will monitor exerciser's heart rate (RS300X, Polar Electro Oy, Kempele, Finland) during the exercise sessions. We will estimate energy expenditure during the exercise sessions by calibrating energy expenditure to heart rate during the VO_{2max} tests done at baseline and at post-intervention time-points. The regression of energy expenditure on heart rate will be calculated for each participant, and energy expenditure during exercise will be estimated from the participant's average heart rate and minutes spent exercising during the training sessions. Additionally, rating of perceived effort will be collected each session [47].

Intensity. Several public health institutions recommend that moderate-intensity physical activity might be beneficial for health in deconditioned persons [43, 45, 46]. Yet, additional benefits have been observed of vigorous vs. moderate-intensity exercise [46]. An intensity of 60% HRres is sufficient to produce clinically significant physiological adaptations in sedentary individuals [43, 45, 46]. The intensity selected for ACTIBATE was 60% HRres for the moderate-intensity group and 80% of HRres for the vigorous-intensity group. The intensity for

the resistance training will be 50% RM and 70% RM for the moderate-intensity and vigorous-intensity group, respectively. 1RM is the maximum amount of weight one can lift in a single repetition for a given exercise. An intensity equivalent to 40-50% of 1RM may be beneficial for improving muscle strength in sedentary persons beginning a resistance training program [48], whereas 60%-70% of 1RM is recommended for novice to intermediate exercisers to improve strength [48]. As the load (i.e. %RM) is not the only variable that influences strength training intensity, we will control variables as speed of movement during both concentric and eccentric phase, recovery time and range of motion. Therefore, we assume that different loads (i.e. 50% RM and 70% RM) will really constitute different training intensities [49-51].

Frequency. Although the physical activity recommendations suggest doing physical activity on most, preferably all days of the week, there was a concern that more than 5 days per week would be an excessive burden and might have undesirable effects on adherence and motivation to the exercise intervention program. Studies on exercise frequency show little differences for 3 or more days per week provided the weekly dose of exercise is attained [46]. Participants in ACTIBATE will be asked to do their weekly exercise dose in 3 to 4 sessions, at their own choice. The length of the training sessions will be adjusted, so that participants attending 3 or 4 days/week have the same weekly dose. Strength training will be performed on 2 of these 3 or 4 days/week, and therefore 1 or 2 sessions per week will consist solely on aerobic exercise. Participants will be advised to not to be out of training for more than 2 consecutive days, and will not be allowed to train <3 days/week or >5 days/week. Participants will be contacted if they miss a scheduled session or if they do not meet the weekly recommendations.

Type of exercise. Activities programmed for the aerobic exercise are cycle ergometer, treadmill and elliptical ergometer. The resistance training program will mainly involve major upper and lower body muscle groups [48].

Training load variation. We are aware that participants might not be immediately capable of exercising at their required volume and intensity dose; therefore, there will be a gradual progression to the assigned exercise dose (see Figure 2).

Aerobic training. Participants of both exercise groups will start with a dose of 75 minutes/week at 60% HRres. The volume will increase 30 minutes/week, so by the 4th week both groups will achieve the 150 minutes/week dose. From this point onwards, the vigorous-intensity group will start a gradual increase of the exercise intensity. For this group, half of the aerobic training time will be performed at a higher intensity, which will be 5% higher than previous week (i.e. 60% HRres the 4th week, 70% HRres the 5th week, etc.), so by the 7th week, both groups will be training at the pre-specified dose.

Strength training. Participants will go through a familiarization period during the first 4 weeks of the study. During this phase, participants will learn the movement patterns that constitute the base of the different exercises (e.g. squat, horizontal pull, vertical push). They will also perform compensatory training such as core stability, flexibility and stabilizers muscles in order to minimize risk of injuries as well as to promote training adherence.

We are aware that both aerobic and strength training load should be incremented as the participants fitness increase. Aerobic training intensity is controlled based on HRres, which means that a higher speed or power has to be selected to achieve a determined percentage of HRres when fitness is increasing. For the strength training, the load equivalent to the participant's 50% RM or 70% RM will be assessed at the beginning of each training phase (i.e. each 5 weeks, see Figure 2). On the other hand, it is well known that different configurations of the strength-training stimulus can elicit different physiological responses (e.g. muscle damage, metabolic stress, etc.) [52]. Taking into account that it is not known which kind of strength training stimulus is better to activate or recruit BAT, ACTIBATE will vary the type of stimuli across the different training phases (Figure 2).

Phases		Familiarization				Phase 1					Phase 2					Phase 3					Phase 4				
Weeks		1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24
Aerobic training	Aerobic training volume (min)	75	105	135	150	120	150	150	150	150	120	150	150	150	150	120	150	150	150	150	120	150	150	120	150
	Intensity (%HRres) moderate-intensity group	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60	60
	Intensity (%HRres) vigorous-intensity group	60	60	60	60	70	75	80	80	80	80	80	80	80	80	80	80	80	80	80	80	80	80	80	80
Strength training	Intensity (%RM) moderate-intensity group	Weight-bearing and elastics bands				Black block					50	50	50	Black block		Black block					50	50	50	50	
	Intensity (%RM) vigorous-intensity group										50	60	70								60	70	70	70	70
	Type of exercises performed	Slow and global movement pattern (light weights)				Exercises localized in major muscle groups					Exercises localized in major muscle groups					Whole body exercises					Exercises localized in mayor muscle groups		Whole body exercises		
	Training stimulus aim	Learning of movement patterns Compensatory training: -Core stability - Flexibility - Stabilizers muscles				Initial adaptations to resistance training Compensatory training: -Core stability - Flexibility - Stabilizers muscles					Session type A: mechanical tension and muscle damage Session type B: Metabolic Stress					Session type A: mechanical tension and muscle damage Session type B: Metabolic Stress					Session type A: mechanical tension and power Session type B: Metabolic Stress				

Figure 2. Training periodization.

Grey blocks represent weeks where the strength training load is reduced in order to learn the proper technique of the exercise used in the next phase. Black blocks represent weeks where strength training mostly consists on repetition maximum (RM) assessment in the different exercises used in the phase. HR: Heart rate reserve.

Exercise training program: Training periodization.

The training program is divided into 5 phases of different duration (Figure 2), starting with a familiarization phase.

Familiarization. This phase corresponds to the first four weeks of the training program. Participants will perform a combined (aerobic plus strength) session 3 times per week, which means that during familiarization phase there will not be “aerobic exclusive” training sessions. A sedentary person is not able to immediately train at the selected doses for ACTIBATE, therefore, the familiarization phase will allow participants to progressively increment the aerobic training volume until the selected dose is achieved (see Figure 2). On the other hand, the strength training will set light loads using elastic bands or weight bearing exercises, that will increased in load and coordinative difficulty as soon as participants are able to perform the exercises with the proper technique. Strength training will be focused on learning the main movement patterns to be used throughout the program (i.e. squat, hinge, bridge, lunge, planks, horizontal and vertical pulls, and horizontal and vertical push) and to improve: a) Core stability; b) Joint stabilizing muscles strength (e.g. rotator cuff, gluteus medius muscle, etc.); c) Balance and standing stability; and d) Flexibility. Strength training on the last week of the familiarization phase will consist on the same exercises performed in phase 1 (see below), but without considering the RM (i.e. light weights), so that participants will learn the proper movement technique.

All training phases after the familiarization period (i.e. phases 1, 2, 3 and 4) have the same structure and duration. Each phase will have 5 weeks, of which the first one will be used to evaluate the strength training load (RM indirect measure) of the exercises used in this phase. Volume of aerobic training will be reduced up to 120 minutes/week during this evaluation week. Selected intensities (50% RM and 70% RM) will be applied during the second, third and fourth week of the phase. Finally, the last week of each phase (i.e. the fifth) will be used to

learn the exercises used in the next phase and its proper technique and, consequently, the load will be reduced.

In every week (except for the evaluation one), participants will perform 2 combined training sessions, and 1-2 aerobic sessions (depending on the selected frequency). Both combined sessions will be different on the structure (so called session type A or B, see Figure 3) or on the exercises performed. During the weeks of the same phase, participants will repeat the same sessions.

□

SESSION TYPE A			SESSION TYPE B			
<i>Warm-up</i>			<i>Warm-up</i>			
MAIN PART	Exercise	Time/sets	MAIN PART	Exercise	Time/sets	
	Aerobic set 1	10 min		Aerobic set 1	10 min	
	Strength exercise I	1set x 10reps		Aerobic set 2	10 min	
	Strength exercise II	1set x 10reps		Strength exercise I	2sets x 10reps	
	Strength exercise III	1set x 10reps		Strength exercise VII	2sets x 10reps	
	Strength exercise IV	1set x 10reps		Strength exercise V	2sets x 10reps	
	Aerobic set 2	10 min		Strength exercise III	2sets x 10reps	
	Strength exercise V	1set x 10reps		Strength exercise II	2sets x 10reps	
	Strength exercise VI	1set x 10reps		Strength exercise VI	2sets x 10reps	
	Strength exercise VII	1set x 10reps		Strength exercise IV	2sets x 10reps	
	Strength exercise VIII	1set x 10reps		Strength exercise VIII	2sets x 10reps	
	Aerobic set 3	10 min		Aerobic set 3	10 min	
	Strength exercise I	1set x 10reps		Aerobic set 4	10 min	
	Strength exercise II	1set x 10reps		<i>Cooling down</i>		
	Strength exercise III	1set x 10reps				
	Strength exercise IV	1set x 10reps				
	Aerobic set 4	10 min				
	Strength exercise V	1set x 10reps				
	Strength exercise VI	1set x 10reps				
	Strength exercise VII	1set x 10reps				
Strength exercise VIII	1set x 10reps					
<i>Cooling down</i>						

Figure 3. Examples of types of a combined training session.

In session type A strength training aims to maximize mechanical tensions and muscle damage, while in session type B strength training aims to maximize metabolic stress. On session type A, the exercise order alternate between upper-body and lower-body, and agonist and antagonist exercises, allowing the maximum recovery possible to the implied muscle groups. On session type B, all upper-body exercises are performed before the lower-body exercises, and all agonist muscle groups exercises are also performed afterward each other.

Phase 1.

Aerobic training volume will be 150 minutes/week (except for the evaluation week, see Figure 2). The vigorous-intensity group will exercise at 70% of HR_{res} half of the aerobic training time (i.e. 60 minutes) in the 5th week of the training program. During the following weeks, the vigorous-intensity group will gradually increase the aerobic target intensity; thus, by the 7th week of the program, this group will train at the selected training intensity.

Strength training will consist on exercises localized in the main muscle groups (e.g. bench press, leg press, lat pull down, etc.) and will include several compensatory exercises (e.g. core stability, stabilizers muscle, etc.) similar to those performed in the familiarization phase. Combined strength sessions (which includes aerobic and strength training) will be a type A session (see figure 3) and will include a total of 4 major muscle groups, 3 core stability and 4 compensatory exercises. Participants will perform a total of 2 sets of 10 repetitions. For the vigorous-intensity group, strength training intensity will also be gradually increased until the selected dose is achieved (See Figure 2).

Phase 2. Combined session will consist on a type A session and a type B session (see Figure 3). Strength training will include similar exercises as those reported in the previous phase, as well as exercises involving several small muscle groups (e.g. arm curl, elbow extension, etc.), performing a total of 8-9 exercises per session. Both, session type A and session type B will include the same strength exercises. Strength exercises will be performed on 2 sets of 10 repetitions each session. On session type B, recovery time will be 1 minute between sets and between different exercises.

Phase 3. It will include a type A session and a type B session (see Figure 3). Both session A and session B will contain the same strength exercises. Strength training will include global exercises, which use simultaneously two or more kinetic chains (e.g. Lunge+horizontal press, hinge+horizontal pull, etc.) and, consequently, constitute a vigorous stimulus to stabilizers

muscles such as core muscles. Strength exercises will be performed on 2 sets of 10 repetitions each session. On session type B, recovery time will be 1 minute between sets and between different exercises.

Phase 4. After the evaluation week, this phase will be divided into two periods of two weeks. It will follow an undulating periodization [53] that will use similar stimulus than that used in phase 2 and 3, but emphasizing power training (i.e. maximum concentric speed, lower eccentric phase time, intra-set recoveries, etc.). The first 2-weeks period will use the same sessions than phase 2 and the second 2-weeks period will contain the phase 3 session types (see above).

Training sessions

The duration of each individual session will depend on the number of visits per week (see above, frequency). Yet exercisers will be advised to evenly distribute the 150 minutes of aerobic physical activity throughout the week, and will be advised not to do aerobic physical activity for longer periods than 60 minutes per session. There was a concern that longer periods of aerobic exercise would be an excessive burden and might have an adverse effect on adherence to the exercise intervention program due to a higher risk of injuries, fatigue and boredom. In order to distribute homogeneously the training time across sessions, aerobic training time will be distributed considering the strength training time, which means that combined training sessions (aerobic training and strength training) will have a lower aerobic volume than the aerobic ones. In every aerobic training session, participants will alternate aerobic exercise with compensatory exercises (e.g. scapular and shoulder mobility and muscles activation, core stability, hip stabilizers muscle, balance exercises, etc.) that constitute a very low training load but might help to avoid boredom during sessions and prevent injuries.

Every combined training session will start with a warm up consistent on 5 minutes of moderate intensity aerobic exercise (i.e. 60%HR_{res}), and a set of mobility and activation exercises (e.g. front plank, monster walker, bridge, pelvic mobility, shoulder rotation, etc.). On aerobic

training sessions, exercisers will only carry out the aerobic part of the warm-up. After warming-up, aerobic training will be performed on series of 10 minutes. Participants will change the ergometer (cycle ergometer, treadmill and elliptical ergometer) between series to avoid overuse of anatomical structures and boredom.

In the combined training sessions type A, strength exercises will be intercalated between aerobic series, mainly to avoid monotony that could result in participants drop out. However in type B sessions, strength training will be performed before the aerobic training, and all aerobic series will be performed one after the other. At the end of every session, participants will perform a cooling-down protocol consistent on 2 minutes of aerobic exercise at a very light intensity and a set of stretching exercises of the main muscle groups involved in the training session.

Usual care group (control). Participants randomly assigned to the usual care (control) group will receive general advices from the exercise-training specialist about the positive effects of physical activity at the start of the study. We will prepare informative pamphlets describing the benefits of physical activity that our group has prepared for the Region of Andalucía (Spain), and that meet with the World Health Organization Guidelines. http://www.juntadeandalucia.es/salud/servicios/contenidos/andaluciaessalud/docs/130/Guia_Recomendaciones_AF.pdf.

Outcomes measures

Primary outcome measures are BAT activity and mass. Secondary outcome variables include REE, MIT, CIT, body temperature regulation and shivering threshold, body composition and cardiovascular disease risk factors. Other variables of interest include cardiorespiratory fitness and muscular strength, dietary habits, physical activity, sleep habits, health-related quality of life and other psychosocial variables, appetite, and demographic characteristics (Table 2).

Table 2. Summary of time point measurements of the primary and secondary outcomes of the study.

Measurements	1- Baseline (week 0)	2- Week 4	3- Week 8	4- Week 12	5- Week 16	6- Week 20	7- Post-int (week 25)
<i>Primary outcome</i>							
BAT activity and mass	x						x
<i>Secondary outcomes</i>							
Resting energy expenditure	x		x		x		x
Meal induced thermogenesis	x						x
Cold-induced thermogenesis	x						x
Shivering threshold	x						x
Body temperature regulation	x						x
Body composition	x		x		x		x
CVD risk factors	x						x
Metabolomics	x						x
Proteomics	x						x
Morphology of tissues	x						x
Dietary assessment	x		x		x		x
Appetite questionnaires	x		x		x		x
Physical activity (monitors)	x		x		x		x
Sleeping habits	x		x		x		x
Cardiorespiratory fitness	x		x		x		x
Muscular strength	x		x		x		x
Health-related quality of life	x						x
Side effects	x	x	x	x	x	x	x

We will also obtain biopsies from abdominal subcutaneous white adipose tissue and skeletal muscle to analyse the expression of genes encoding proteins involved in the thermogenic machinery.

The baseline assessments are divided into 5 days:

Day 1: During the first session, each participant will go through a medical examination by the sports medicine staff from the *Centro Andaluz de Medicina del Deporte (CAMD)*. Resting electrocardiogram and blood pressure will be measured. Thereafter, participants will perform muscle strength tests (see below). Lastly, a maximum exercise test will be conducted on a treadmill (h/p cosmos, Italy).

Day 2: During the second assessment day, participants will arrive at 8:00 to the laboratory, where they will rest for 60 minutes during which time heart rate variability will be measured the first 15 minutes (Polar RS800X, Polar Electro Öy, Kempele, Finland), and REE will be measured the last 30 minutes by indirect calorimetry. Afterwards, they will have a standard

breakfast (50% REE, T-Diet, Vegenat[®]: 16.Kcal/ml; 47% CHO, 30% FAT; 15% PRO, 3% FIBER) and energy expenditure will be measured for other 3:30 hours to determine MIT [54]. Appetite measures [visual analogue scales, (VAS)], thermal sensation and thermal comfort (VAS), and skin temperature will be monitored during and after the MIT. Lastly, body composition will be determined by Dual Energy X-ray Absorptiometry (DXA) scan (HOLOGIC, QDR 4500W).

Day 3: We will determine shivering threshold (Polar Products Inc, Ohio, USA) and CIT.

Day 4: We will conduct PET/CT scans (Siemens Biograph 16 PET/CT, Siemens, Germany), and we will collect blood sample to determine energy metabolism-related biomarkers changes in response to cold, at the *Hospital Universitario Virgen de las Nieves (Granada, Spain)*.

Day 5: Fasting blood measurements and muscle and subcutaneous adipose tissue biopsies.

In a separate day, participants will fill in at home several web-based questionnaires regarding dietary habits, quality of life, sleeping habits, etc. Physical activity and sleep quality will be assessed through accelerometry during a week, and dietary intake will be measured by three 24h recalls.

Primary outcome (objective 1): To quantify the effect of exercise intensities on BAT activity and mass, we will determine the maximal and mean standardized uptake values. We will do a PET/CT scan (Siemens Biograph 16 PET/CT, Siemens, Germany) to each participant at the cervical and supraclavicular level until mediastinum, after an individualized cooling protocol [55]. Participants will rest in a room at 18-19°C for an hour wearing a temperature controlled water circulation cooling vest (Polar Products Inc., Ohio, USA) at a temperature 3°C above the individually measured shivering threshold (see below). One hour later, we will raise 1°C the water temperature, and we will inject 74MBq (2 mCi) of ¹⁸F-fluorodeoxyglucose (¹⁸F-FDG) tracer and will rest for another hour with the cooling chest vest at the same room temperature. If the participant reports shivering, the water temperature will increase 1°C until shivering

stops. During all the process the skin temperature will be monitored, as well as the systolic and diastolic blood pressure. Two nuclear medicine physicians will independently interpret the images from the PET/CT. Additionally, immediately before the cold exposure (thermoneutral conditions) and after one hour of cold exposure (before the tracer injection) we will collect a blood sample to determine markers of energy metabolism such as glucose, insulin, and non-esterified fatty acids.

Secondary outcomes

To investigate the metabolic consequences of activating BAT we will measure:

REE. It will be measured by indirect calorimetry (CCM express®, Medgraphics, Saint Paul, MN USA). REE will be calculated from O₂ and CO₂ volumes, as well as from urine excretion nitrogen values.

MIT. It will be determined as the increase in energy expenditure above the REE divided by the energy content of a standardized meal (50% REE, T-Diet Vegenat).

CIT. We will measure energy expenditure with indirect calorimetry during cold exposure of the aforementioned cooling protocol.

All calorimetric measures will follow the accepted standard to assure the validity of the tests [56].

Shivering threshold. To determine the shivering threshold, participants will wear a temperature controlled water circulation cooling vest (Polar Products Inc., Ohio, USA). The cooling vest covers the clavicular region of the participant, as well as the chest and abdominals. During the cooling protocol, participants will stay in a room at a temperature of 18-19 °C in a semi-supine position. The cooling starts at a water vest temperature of 21°C and water will be gradually decreased until shivering occurs [57]. We will determine shivering both visually and by asking the participants if they are experiencing shivering.

Body temperature regulation. We will monitor skin temperature during (i) cooling protocols previous to the PET/TC; (ii) shivering threshold protocol; (iii) MIT test; and (iv) exercise test. Participants will wear a set of iButtons (DS 1922L, Thermochron, iButtons®; resolution: 0.0625°C, accuracy: 0.5°C; Maxim, Dallas, USA) covering the standard skin points following ISO 9886-2004 [58] defined locations to measure skin body temperature, proximal, distal and skin temperature [59]. Peripheral vasoconstriction will be estimated through the gradient of two iButtons (forearm and fingertip) [60]. Moreover, subjective cold perception will be assessed by VAS. We will also monitor supraclavicular skin temperature since it seems to have predictive value for BAT detection in adult humans [57].

White adipose tissue and muscle biopsies. To analyse the expression of genes encoding proteins involved in the thermogenic machinery, mitochondriogenesis and Brite brown adipocytes, we will obtain biopsies from abdominal subcutaneous white adipose tissue at the umbilical level to quantify whether there exists browning in white adipose tissue depots, expression of UCP-1 and other potential targets involved in the browning process; and in skeletal muscle at *vastus lateralis* to quantify the expression of genes encoding proteins potentially involved in human thermogenesis. An expert surgeon will perform the biopsies following the Bergström's technique. Immunohistochemical staining and quantitative RT-PCR will be used to perform these analyses.

Plasma samples and biochemical analysis. We will also measure a full set of cardiovascular risk factors including fasting plasma lipids and lipoproteins, glucose and insulin, myeloperoxidase, e-selectin, vessel cell adhesive protein 1, and immunoglobulin cell adhesive protein, cytokines (e.g tumor necrosis factor alpha, IL-6 and monocyte chemoattractant protein 1) adipokines (e.g leptin, adiponectin, resistin and active plasminogen activation inhibitor factor 1) as well as irisin and soluble (extracellular) FNDC5, orexin A and B neuropeptides, atrial natriuretic peptides, bone morphogenetic protein 7, fibroblast growth factor 21, transforming

growth factor- β , and β -aminoisobutyric acid, among others. ELISA kits, western blots and High Performance Liquid Chromatography- Mass Spectroscopy will be used to perform these analyses.

Cardiorespiratory fitness. It will be determined using a maximum treadmill exercise test (h/p cosmos, Italy) following the modified Balke protocol [61], which has been extensively used and validated [62-64]. The criteria for achieving maximum oxygen consumption (VO_{2max}) will be: respiratory exchange ratio ≥ 1.1 , a plateau in VO_2 (change of <100 ml/min in the last three consecutive 10 seconds stage), and a heart rate within 10 beats/min of the age-predicted maximal heart rate ($209-0.73*age$). Participants that do not achieve these criteria will be classified as having reached the VO_{2peak} . Subsequent analysis will account for this factor. The exercise electrocardiogram will be monitored continuously.

Muscular strength. We will determine 1RM of the bench press and leg press, using the resistance weight machines (Keiser $\text{\textcircled{R}}$) that will be used in the exercise training sessions. We will also measure handgrip strength [65].

Appetite. To examine whether changes in energy expenditure (REE, MIT, CIT) during and post-exercise intervention lead to increased appetite and therefore to an increased energy intake, we will assess (i) appetite through questionnaires (Three-Factor Eating Questionnaire, Binge Eating Scale, Control of Eating Questionnaire), and VAS; (ii) **Dietary intake.** It will be estimated by means of a previously validated food frequency questionnaire and 3 non-consecutive 24h dietary recalls (one weekend day and two non-weekend days). Dietary and nutritional assessment will be analysed using a database for the composition of Spanish foods. We will remind participants not to change their dietary habits.

Body composition. We will also measure fat mass, lean body mass, abdominal adipose tissue and bone mineral density by conducting a DXA scan (HOLOGIC, QDR 4500W). We will also measure waist circumference.

Physical activity. We will monitor unstructured daily physical activity to assess potential changes in non-supervised physical activity, and to assess non-exercise activity thermogenesis. All participants will wear two activity monitors, one on the wrist and another on the hip (ActiSleep, Actigraph, Pensacola, Florida, USA) for 7 consecutive days for 24 hours to record physical activity intensity levels and patterns, as well as sleeping habits. Participants will also complete a diary log.

Adverse effects. We will record adverse effects or health problems attributable to the testing sessions or intervention sessions, including muscle pain, fatigue, and general aches and pains by self-report during the study period. An independent researcher will be in charge of auditing all assessment staff to record all these events in the participants over the study period.

Room and outdoor temperature. The temperature of the exercise room as well as outdoor will be recorded all through the study from the Spanish Weather Service (*Agencia Española de Meteorología*, www.aemet.es/es/portada). Participants will exercise at 20-22°C according to the American College of Sports Medicine [66].

Potential impact of ACTIBATE

This project contributes to one of the social challenges set up by the Horizon 2020 as priority: Health, demographic change and well-being. It will improve our knowledge to combat chronic diseases as obesity and diabetes. Currently, there are no simple and non-surgical treatment strategies for treating type obesity, type 2 diabetes, and atherogenic development. While physical exercise seems to play a key role in the prevention and treatment of obesity and insulin resistance, the mechanisms that may mediate these effects are currently unknown. ACTIBATE will quantify, for the first time in humans, the effect of physical exercise on BAT activity and mass measured by PET/CT (primary outcome). Moreover, ACTIBATE will obtain biopsies from white adipose tissue and skeletal muscle tissues to analyse the expression of genes encoding proteins involved in the thermogenic machinery. We will also investigate the

metabolic consequences of activating BAT by measuring REE, MIT, CIT, body temperature regulation and shivering threshold, and cardiovascular disease risk factors before and after the exercise intervention. This study integrates three different disciplines, namely exercise physiology, nutritional biochemistry, and nuclear medicine, to investigate whether exercise is able to (i) activate and recruit BAT, (ii) identify new BAT precursors and (iii) induce the specific gene program to favour white-to-brown adipocyte transformation in humans. The exercise intervention may potentially translate to a modest (around 5%) but “chronic” increase of REE. Moreover, even if BAT recruitment would not produce enough energy expenditure increase to induce a negative energy balance, it could potentially exert metabolic benefits. Findings from the ACTIBATE will have significant implications for our understanding of exercise and its protective effects against the development of type 2 diabetes, obesity and related metabolic diseases.

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