Large muscles are beneficial but not required for improving thermogenic capacity in small birds

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It is generally assumed that small birds improve their shivering heat production capacity by developing the size of their pectoralis muscles. However, some studies have reported an enhancement of thermogenic capacity in the absence of muscle mass variation between seasons or thermal treatments. We tested the hypothesis that an increase in muscle mass is not a prerequisite for improving avian thermogenic capacity. We measured basal (BMR) and summit (M\text{sum}) metabolic rates of black capped chickadees (Poecile atricapillus) acclimated to thermoneutral (27 °C) and cold (−10 °C) temperatures and obtained body composition data from dissections. Cold acclimated birds consumed 44% more food, and had 5% and 20% higher BMR and M\text{sum}, respectively, compared to individuals kept at thermoneutrality. However, lean dry pectoralis and total muscle mass did not differ between treatments, confirming that the improvement of thermogenic capacity did not require an increase in skeletal muscle mass. Nevertheless, within temperature treatments, M\text{sum} was positively correlated with the mass of all measured muscles, including the pectoralis. Therefore, for a given acclimation temperature individuals with large muscles do benefit from muscle size in term of heat production but improving thermogenic capacity during cold acclimation likely requires an upregulation of cell functions.

For small avian species wintering at high latitudes, winter acclimatization is mainly a physiological phenomenon where cold hardiness is improved as temperature decline from fall to peak of winter. This improved capacity is typically associated with increases in basal (BMR) and summit (M\text{sum}) metabolic rates, which are respectively thought to reflect physiological maintenance costs and cold endurance.

The seasonal elevation in BMR is often interpreted as resulting from an increase in daily food consumption requiring larger digestive and excretory organs (e.g. liver, gizzard, intestine), in turn leading to higher maintenance costs. However, as the influence of body composition on BMR is context-specific and can be affected by tissue metabolic intensity, this scenario may not be generalizable. For example, in cases where acclimatization also leads to considerable increases in skeletal muscle size, the influence of digestive and excretory organs on BMR can be overshadowed by the amount of muscle tissues consuming energy during measurements. In contrast, since M\text{sum} is a measure of maximal shivering heat production, the influence of skeletal muscle size, particularly the flight muscles and heart size, on thermogenic capacity appears much more consistent. In several small free-living wintering species, elevated winter M\text{sum} is indeed associated with seasonally larger pectoralis muscles. The mass of skeletal muscles and heart has also been found to correlate significantly and positively with M\text{sum} several times.

Despite the seasonal changes in muscles size and M\text{sum} observed in the wild, a small number of studies, although they were not designed to investigate this specific phenomenon, reported improvements of thermogenic capacity in controlled conditions independently from changes in muscles size. This phenomenon was found to occur even in species known to increase pectoralis muscle mass in winter. For example, dark-eyed juncos (Junco hyemalis) are known to increase both their M\text{sum} and the size of their pectoralis muscles in winter relative to summer but recently, Swanson et al. conducted an experiment with captive juncos and found a 16–19% higher M\text{sum} in cold-acclimated (3 °C) birds relative to individuals exposed to a warm treatment (24 °C) with no difference in pectoralis muscle mass. Similarly, Barceló et al. documented a 19% higher M\text{sum} in captive white-throated sparrows (Zonotrichia albicollis) acclimated to –8 °C compared to individuals maintained in a warm condition (24 °C) with no difference in pectoralis muscle mass.

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at thermoneutrality (28 °C). In this particular case, although the expected positive correlation between heart and muscles mass and M\textsubscript{sum} was found in cold acclimated birds, there was no significant difference in the mass of pectoral or other skeletal muscles between thermal treatments.

Since shivering does not produce external work, and thus most of the chemical energy consumed during contraction is released as heat\textsuperscript{36}, there is no doubt that large muscles should produce more heat, for a given level of shivering, compared to small muscles. Consequently, in a given dataset if muscles mass and M\textsubscript{sum} cover a range of variation sufficiently wide, which is often the case with intersessional studies\textsuperscript{26,10,31,35}, positive correlations between muscle mass and M\textsubscript{sum} should be detectable across or within seasons. However, the experimental evidence presented above suggest that developing larger muscles may not be an obligate prerequisite for improving individual thermogenic capacity, even in species known to increase the size of their muscles during cold winters.

To test this hypothesis, we conducted an experimental study with captive black-capped chickadees (\textit{P. atricapillus}). Chickadees are small (11 g) non-migratory passerines that typically express elevated M\textsubscript{sum} in winter relative to summer\textsuperscript{4,26,21}. This improvement of thermogenic capacity parallels or statistically correlates with the development of larger muscles and heart\textsuperscript{26,31,42}. However, at least one case of chickadees going through a cold winter without significant changes in pectoral muscles size has been documented\textsuperscript{44}. This suggests that in experimental conditions, where only temperature is manipulated, this species could also show improvement of thermogenic capacity independently from skeletal muscle mass variation. We therefore exposed birds to two thermal treatments (−10 °C and 27 °C), with the expectation that cold acclimated birds, would show a higher M\textsubscript{sum} than those maintained at thermoneutrality but no difference in mean mass of pectorals and other skeletal muscles. We nevertheless expected a correlation between muscle mass and M\textsubscript{sum} across or within treatments as individuals with larger muscles could still benefit from the mass of these tissues in terms of maximal shivering heat production, independently from their acclimation temperature. We also measured BMR in these birds to document maintenance costs. In this particular case, we expected that BMR variation across treatments would correlate with the mass of digestive and excretory organs if there was no major difference in muscle mass between treatments\textsuperscript{49}. In contrast, we expected BMR to correlate with the mass of skeletal muscles if cold acclimated birds enlarged the size of these organs\textsuperscript{26}.

Material and Methods

Birds collection and acclimation. From January to April 2015, we captured 49 black capped chickadees using mist nets at two sites, the Forêt d’Enseignement et de Recherche Maçêts, (48°19N, 68°30W) and lac à l’Anguille (48°25N, 68°25W), both in eastern Québec, Canada. These birds were brought into captivity at the avian facilities of the Université du Québec à Rimouski. Birds were held in individual cages (39 × 43 × 31 cm) and exposed to a constant photoperiod (10 L:14D) for the remainder of the experiment. Birds consumed a diet of living mealworms and freshly-thawed crickets (0.20 g and 0.30 g per day, respectively), sunflower seeds, Mazuri small birds maintenance diet (MAZURI® exotic animal nutrition, USA) and water, which were available ad libitum. The birds also received vitamin supplements, daily (Electolytes plus, Vetoquinol N.-A.INC, QC, Canada) and once per week (Poly-tonine A® complex, Vetoquinol N.-A.INC, QC, Canada) and once per week (Poly-tonine A® complex, Vetoquinol N.-A.INC, QC, Canada) in their water. Our experimental groups were formed of 24 individuals maintained at −10 °C (cold) and 25 individuals maintained at 27 °C (thermoneutral zone of this species\textsuperscript{4,44}). Birds were acclimated to these conditions for a minimum of 39 days (mean = 61.5, max = 84) after which we measured average daily food consumption over 6 days (same diet but excluding Mazuri) by subtracting the mass of food left in food trays in the morning from what had been offered the day before at the same time. Following the 6 days of food intake measurements, we proceeded with metabolic rate trials (see below).

All bird manipulations have respected the Canadian Council on Animal Care (CCAC) guidelines and were approved by the animal care committee of the Université du Québec à Rimouski (CPA-60-15-160). They also have been conducted under scientific and banding permits from Environment Canada–Canadian Wildlife Service.

BMR and M\textsubscript{sum} measurement. Because tissues collected on birds were also analyzed in another experiment that required fresh samples (results not shown, Milbergue et al., in prep), our measurements sequence for metabolic rate was limited to recording BMR and M\textsubscript{sum} on 8 birds per week, until all birds had been measured (49 days). Each day of respirometry trial involved measurement on four birds (2 from each treatment) and followed the protocols described in details by Lewden \textit{et al.}\textsuperscript{46} and Petit \textit{et al.}\textsuperscript{8,26} where the animals VO\textsubscript{2} were measured using FoxBox oxygen analyzers (Sable Systems, Las Vegas, NV, USA). Each M\textsubscript{sum} trials were conducted on two randomly chosen birds from a same temperature treatment and began approximately at 9:00 and at 12:30 (alternating treatments between measures). Trials began by weighing the birds (0.00 g, Scout Pro, Ohaus, NJ, USA) and placing each of them individually in a stainless steel metabolic chamber (volume = 1350 ml). The birds then received air during 20 min before being exposed to helox gas (21% oxygen, 79% helium) using a flow rate of 900 ml min\textsuperscript{−1} controlled by mass flow controllers (Omega, FMA 5400/5500, QC, Canada) calibrated with a Bubble-O-Meter (Dublin, OH, USA). We used a sliding cold exposure protocol\textsuperscript{36}, where ambient temperature was first set to either 0 °C (cold group) or 10 °C (thermoneutral group) and then ramped down by 3 °C every 20 min. Trials ended when birds became hypothermic, which was easily identifiable in real time as a steady decline in oxygen consumption for several minutes. Body temperature was immediately measured after taking birds out of their chamber using a thermocouple reader (NIST-traceable Omega model HH-25KC, QC, Canada) and a copper constantan thermocouple inserted into the cloacae, approximately 10 mm deep. Only data from birds showing a body temperature after trials lower or equal to 38.5 °C\textsuperscript{46,48} were used in the analyses. This removed five M\textsubscript{sum} measurements from our sample. Body mass was again recorded at the end of trial and average body mass was used in statistical analyses on M\textsubscript{sum}.
After $M_{\text{sum}}$, birds were brought back to their cage and had access to food and water until BMR measurement, starting at around 19:00. BMR trials were done on all 4 birds at 30 °C (thermoneutral zone) in chambers that received 500 ml min$^{-1}$ of dry, CO$_2$ free air. Trials ended the following morning (at approximately 7:30). As for $M_{\text{sum}}$, body mass was measured prior to and after BMR measurement and the average was used in statistical analyses. Birds were then returned to their cage.

Metabolic rates were calculated with the EXPEDATA software, v.1.8.4 (Sable Systems, Las Vegas, NV, USA) using the equation 10.1 of Lighton. $M_{\text{sum}}$ and BMR were calculated from the highest and lowest averaged 10 min of VO$_2$. Because birds use lipids as metabolic fuel during shivering and the duration of BMR trials (> 720 min) insured that birds were post-absorptive at time of BMR measurement, we estimated heat production in Watts assuming an energy equivalent for lipid oxidation of 19.8 kJ l$^{-1}$ O$_2$. Five individuals died of unknown cause during the experiment, leaving a final sample size of $n = 20$ (−10 °C) and 18 (27 °C) for BMR and $n = 18$ (−10 °C) and 19 (27 °C) for $M_{\text{sum}}$.

**Organ collection.** Birds were euthanized by decapitation in the 2 to 5 days following their respirometry trial (delay caused by measurements conducted on tissues in parallel to this experiment, Milbergue et al., in prep). The right and left pectoralis muscles, heart, liver, empty intestine, pancreas and gizzard were removed within minutes of the birds death and weighed (0.0001 g) with a precision balance (Cole-Parmer Symmetry, PA-Series, Canada). These organs were placed in Eppendorf tubes and immersed in liquid nitrogen before being transferred to a −80 °C freezer. Carcasses were preserved at −20 °C until we completed dissections. This was done by removing and weighing the brain, lungs, kidneys, skin (feathers removed) and upper right and left leg muscles, considered as a single organ and including bones. The remaining carcasses were therefore composed mainly of skeletal muscles and bones. Organs were then freeze-dried (FreeZone 2.5, Labconco, Kansas city, KS, USA) for 2 days to obtain constant dry mass of tissues. Adipose tissues have low metabolic activity and can bias analyses on relationships between mass or body composition and metabolic rates when birds contain differing amounts of fat. We therefore extracted lipids from these samples with a Soxhlet apparatus using petroleum ether to obtain final lean dry mass of organs.

It should be noted here that we do not have lean dry mass data for the heart as the entire organ was needed for tissue analyses (Milbergue et al., in prep). Wet mass is therefore presented for this organ. For pectoralis muscles and liver, since we used subsamples for tissue analyses and processed the remaining tissue as the other organs (freeze-drying and fat extraction), we recalculated lean dry mass of these organs in proportion to the original wet mass of the complete organ. In this experiment, we originally planned on obtaining ash-free lean dry mass for leg muscles and the remaining carcass but a technical problem during the burning of samples in a furnace led to the loss of a large number of samples. We therefore cannot present ash-free data.

**Statistical analysis.** Our analyses first tested whether birds differed between treatments before the temperature change. We thus used one-way ANOVAs to test for a treatment effect (cold or thermoneutral) on furcular fat score (estimated according to Gosler), structural body size and body mass measured prior to group formation. Structural body size was calculated as the first principle component (PC) from a principal component analysis combining variation in length measurements of head plus beak, tarsus, wing and tail.

To determine how thermal environments might have influenced body composition after acclimation, we ran ANCOVA models testing for the effect of thermal treatment on organ lean dry mass. Since structurally larger birds might also have larger organs, we included body size as a covariate in these models. We used the same approach to determine the influence of thermal treatments on BMR and $M_{\text{sum}}$. Models included the effect of time since capture to consider a potential influence of captivity duration, but this last variable was not significant and is therefore not considered further. The models were first run on whole BMR and $M_{\text{sum}}$. We then included structural body size or body mass as covariate but the size effect was not significant in any models thus this effect is not presented here.

To determine the influence of body composition on metabolic performance, analyses are typically based on stepwise regressions or a model selection approach where the influence of all body constituents on BMR and $M_{\text{sum}}$ are compared and ranked in order of significance and importance of their effect (e.g. 25,28). However, results from these analyses depend on the variables included in models and missing variables can influence results. In the present case, we could not include lean dry heart mass in our analyses. However, although this organ typically represents only 1% of total body mass, the heart has been shown to significantly contribute to variation in both BMR and $M_{\text{sum}}$. We therefore chose a simpler approach for our analyses. We conducted separate ANCOVA models including thermal treatment, lean dry mass of the organ (fresh mass for heart) and their interaction. These models were then ranked according to the Bayesian information criterion (BIC).

Analyses testing for the effect of muscle tissues on metabolic rates were first conducted considering muscle groups separately. Models thus included either pectoralis muscles, leg muscles (including bones) or carcass (including bones) as independent variables. As leg muscles and carcass mass included bones and bone mass should closely correlate with structural body size, we also included body size as an additional covariate in these models to control for bone mass. Then, we combined pectoralis muscles, legs muscles and carcass to generate a “total muscle” variable and used total muscle as our independent variable in the model. Here again, structural size was included as a covariate to control for bones mass. In all of these cases, however, the effect of structural body size was never found to be significant, likely because bones mass (measured as ash) only represents a small proportion of lean dry body mass in chickadees (27% of carcass mass, including all bones, based on data from 26). This effect is therefore not included in the models presented here.
Results

Treatment effect on body composition. Birds from both groups did not differ prior to treatment. There was no significant effect of treatment on furcular fat score ($F_{1,24} = 2.4, P = 0.1$), structural body size ($F_{1,26} = 1.2, P = 0.3$) or body mass ($F_{1,27} = 0.97, P = 0.3$). Groups did not differ in sex ratio either ($\chi^2 = 0.19, P = 0.66$, sex determined during dissection). After 28 days of acclimation, differences were detected (Table 1). At the end of acclimation, cold-acclimated birds were eating 43.5% more food but nevertheless had, at the time of dissection, 27–30% less fat than individuals maintained at thermoneutrality (Table 1). Despite this difference in fat content, post-acclimation body mass did not differ significantly between treatments (Table 1). Among organs, only the heart (fresh mass), lungs, pancreas and leg muscles differed between experimental temperatures (Table 1). Cold-acclimated birds had a 10.7% and a 8.0% smaller lungs and legs muscles but had a 14.3% and a 95.8% larger heart and pancreas, respectively. No other organs differed between treatments, including pectoralis muscles, carcass, digestive and excretory organs. Total lean dry muscles mass differed by less than 1% between treatments (Table 1).

Influence of temperature and body composition on metabolic performance. Cold-acclimated birds had a BMR 4.5% higher on average than individuals kept at thermoneutrality, but the temperature effect depended on body mass (interaction body mass*treatment, Table 2). Indeed, although the influence of body mass on BMR was clear at thermoneutrality (independent regression $R^2 = 0.61, n = 19, P < 0.0001$), this effect appeared uncoupled at $-10\,^\circ$C (independent regression: $P = 0.99$, Fig. 1a). Therefore, most birds kept in the cold had a BMR as high as the heaviest birds kept at thermoneutrality (Fig. 1a). Ranking independent ANCOVA models for relationship between organ mass and BMR revealed a clear influence of skeletal muscle mass on BMR (Table 3).

Table 1. Least square means (±s.e.m) and differences between cold ($-10\,^\circ$C: C) and thermoneutral (27 \,^\circ$C: T) treatments for body composition variables in black-capped chickadees. Units are in grams except for food intake (g/day). *Includes bone mass and controls for structural body size (see text for details).

| Variable | Cold | Thermoneutral | F (df) | P | % difference (C relative to T) |
|----------|------|---------------|--------|---|-----------------------------|
| Food intake | 3.86 ± 0.09 | 2.69 ± 0.08 | 96.4 (1,45) | <0.0001 | 43.5 |
| Mass and fat | | | | | |
| Body mass | 11.96 ± 0.16 | 12.25 ± 0.16 | 1.5 (1,47) | 0.2 | −2.4 |
| Total organ fat mass | 0.88 ± 0.07 | 1.25 ± 0.06 | 16.1 (1,35) | <0.001 | −29.6 |
| Furcular fat mass | 0.11 ± 0.01 | 0.15 ± 0.01 | 5.7 (1,35) | <0.05 | −26.7 |
| Muscles | | | | | |
| LD pectoralis | 0.41 ± 0.01 | 0.39 ± 0.01 | 0.97 (1,34) | 0.33 | 5.1 |
| LD legs* | 0.23 ± 0.005 | 0.25 ± 0.004 | 9.2 (1,33) | <0.01 | −8.0 |
| LD carcass* | 1.21 ± 0.02 | 1.24 ± 0.02 | 0.83 (1,33) | 0.37 | 2.4 |
| LD total muscles* | 1.86 ± 0.04 | 1.85 ± 0.04 | 0.0005 (1,33) | 0.98 | 0.5 |
| Cardio pulmonary | | | | | |
| Heart | 0.16 ± 0.004 | 0.14 ± 0.004 | 12.0 (1,34) | <0.001 | 14.3 |
| LD lungs | 0.025 ± 0.001 | 0.028 ± 0.001 | 6.4 (1,35) | <0.05 | −10.7 |
| Digestive and excretory | | | | | |
| LD gizzard | 0.087 ± 0.005 | 0.079 ± 0.005 | 1.35 (1,35) | 0.25 | 10.1 |
| LD intestine | 0.039 ± 0.002 | 0.036 ± 0.002 | 1.6 (1,34) | 0.22 | 8.3 |
| LD liver | 0.078 ± 0.004 | 0.066 ± 0.004 | 2.0 (1,25) | 0.17 | 18.2 |
| LD pancreas | 0.0094 ± 0.0007 | 0.0048 ± 0.0007 | 23.6 (1,32) | <0.0001 | 95.8 |
| LD kidneys | 0.030 ± 0.001 | 0.029 ± 0.001 | 0.21 (1,35) | 0.65 | 3.4 |
| Other | | | | | |
| LD skin* | 0.11 ± 0.003 | 0.11 ± 0.003 | 0.01 (1,33) | 0.91 | 0.0 |
| LD brain | 0.10 ± 0.001 | 0.11 ± 0.001 | 0.78 (1,35) | 0.38 | −9.1 |
| Total LD body mass* | 2.33 ± 0.05 | 2.30 ± 0.04 | 0.18 (1,33) | 0.67 | 1.3 |

Table 2. Effects of thermal treatments, body mass on BMR and $M_{\text{sum}}$ and least square means (±s.e.m) per treatment.

| Treatment | F (df) | P | Body mass | Interaction | Cold | Thermoneutral | % difference |
|-----------|--------|---|-----------|-------------|------|---------------|-------------|
| BMR | 7.15 (1,33) | <0.05 | 0.0002 (1,33) | 0.99 | 5.65 (1,33) | <0.05 | 0.23 ± 0.005 | 0.22 ± 0.004 | 4.5 |
| $M_{\text{sum}}$ | 36.8 (1,36) | <0.0001 | 16.1 (1,36) | <0.001 | — | — | 1.65 ± 0.03 | 1.38 ± 0.03 | 19.6 |
The model including total muscle mass (Fig. 1b) ranked first followed by the models including skin, carcass, leg and pectoralis muscle (Fig. 1c). Digestive, cardio-pulmonary and excretory organs all ranked after these organs and thus apparently had less influence on BMR variation (Table 2).

Msum was 19.6% higher in cold-acclimated birds when considering the significant effect of body mass (Table 2, Fig. 2a). As for BMR, muscles had a prominent influence on Msum variation. Total muscles (Fig. 2b), pectoralis muscles (Fig. 2c), leg muscles and carcass were all positively correlated with Msum and ranked first based on BIC values for independent ANCOVA models (Table 4).

### Discussion

In this experiment, we expected higher thermogenic capacity in cold-acclimated birds relative to individuals kept at thermoneutrality and predicted that this difference would not result from larger skeletal muscles in the cold. Our data support this hypothesis as cold-acclimated individuals had a Msum 20% higher but had smaller leg muscles and did not develop larger pectoralis, carcass or total muscles than birds kept at 27 °C. We also expected

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**Figure 1.** Relationships between BMR and body mass or lean dry mass of skeletal muscles in black-capped chickadees: (a) body mass, (b) total lean dry muscle mass (including bones), (c) lean dry pectoralis muscle mass (filled circles: 27 °C, open circles: −10 °C).
that variation in BMR would correlate with digestive and excretory organs if there were no major differences in muscle mass between treatments, or with skeletal muscles if their development was part of the response to cold. This hypothesis was only partially supported since skeletal muscles did correlate positively with BMR across treatment despite a lack of increase in muscle mass in the cold.

Birds acclimated to −10 °C consumed on average 44% more food per day during the last 6 days of acclimation than individuals kept at 27 °C. They also had 5% higher maintenance costs, based on average BMR, and 20% higher thermogenic capacity, based on average $M_{\text{sum}}$. Therefore, the cold treatment was associated with considerable demands for thermoregulation and a consequent physiological response. Nevertheless, body mass did not differ between treatments. In fact, although we found the expected larger heart (based on wet mass) and larger pancreases in cold-acclimated birds, these individuals carried 27–30% less body fat and had smaller lungs and leg muscles than birds acclimated to thermoneutral conditions. The other components of lean body mass did not differ significantly between temperatures although most had higher mean values in cold-acclimated birds (Table 1).

Table 3. Correlations between BMR and body composition. Results are from final ANCOVA models, including lean dry mass of organ and treatment as variables. *Includes bone mass (see text for details).

| Organ       | Treatment | Interaction | Adjusted $R^2$ | BIC  | ΔBIC |
|-------------|-----------|-------------|----------------|------|------|
| Total muscles | 9.8 (1.31) | <0.01       | 0.33           | −163.4 | −     |
| Skin        | 9.0 (1.31) | <0.01       | 0.32           | −162.7 | −0.7 |
| Carcass     | 8.9 (1.31) | <0.01       | 0.32           | −162.6 | −0.8 |
| Legs        | 6.4 (1.31) | <0.05       | 0.28           | −160.5 | −3.0 |
| Pectoralis  | 4.9 (1.31) | <0.05       | 0.24           | −159.0 | −4.4 |
| Lungs       | 1.8 (1.30) | 0.19        | 0.26           | −157.2 | −6.2 |
| Brain       | 2.6 (1.31) | 0.11        | 0.19           | −156.8 | −6.6 |
| Kidneys     | 2.3 (1.31) | 0.14        | 0.19           | −156.5 | −6.9 |
| Heart (wet) | 0.99 (1.30)| 0.33       | 0.24           | −156.3 | −7.1 |
| Intestine   | 6.1 (1.31) | <0.05       | 0.17           | −155.6 | −7.8 |
| Gizzard     | 0.74 (1.31)| 0.4        | 0.15           | −154.9 | −8.6 |
| Pancreas    | 1.2 (1.28) | 0.3        | 0.15           | −138.6 | −24.8 |
| Liver       | 0.12 (1.22)| 0.7        | −0.04          | −102.9 | −60.6 |

In the observed scenario where temperature did not influence skeletal muscle mass, we expected a significant effect of digestive and excretory organs on BMR at lower temperatures. Instead, total mass of muscles across treatments was positively related to maintenance costs. Despite the elevated food intake in the cold, none of the digestive and excretory organs, except pancreas, responded significantly to temperature and all of these organs ranked after skeletal muscles for their importance on explaining BMR. Skeletal muscles represent 73% of ash-free lean dry body mass in free-living wintering chickadees (Petit and Vézina, unpublished data) and pectoralis muscles alone represented 17% of total lean dry body mass in "our birds (Table 1)". Therefore, without major changes in digestive and excretory organs, the energy consumed by resting skeletal muscles in birds under standard BMR conditions likely overshadowed the influence of other metabolically active organs such as the gut, liver or kidneys. This correlation between pectoralis or skeletal muscles and BMR has previously been observed in other avian species, including free-living black capped chickadees.26
With regards to the relationship between skeletal muscle mass and BMR, an important point to consider is that it was only apparent in the thermoneutral group (see Fig. 1b,c). Independent regression models for total skeletal muscles, pectoralis, leg muscles and carcass were all significant at 27 °C ($R^2 = 0.35–0.54$, all $P < 0.01$) but the same analyses for birds acclimated to $-10$ °C yielded no significant relationships ($P > 0.6$ in all cases). As stated earlier, and since most of lean body mass was made of skeletal muscles, this uncoupling in cold acclimated birds is likely resulting from changes in metabolic intensity taking place at the cellular level.

For a given body mass, cold-acclimated black capped chickadees had 20% higher thermogenic capacity than birds maintained at thermoneutrality (Fig. 2a). That difference did not result from larger skeletal muscles since these organs did not differ between groups (<1% difference in total muscle mass) or were smaller (legs) in cold-acclimated birds (Table 1). The combined mass of the heart and lungs has previously been shown to correlate with $M_{\text{sum}}$ variation across seasons in our chickadee population\textsuperscript{26}. Recent evidence also suggests that cardiovascular functions could play a significant role in thermogenic capacity as larger hearts\textsuperscript{18,25,32,33,34,35,36,37,38,39,40,41} and upregulated

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**Figure 2.** Relationships between $M_{\text{sum}}$ and body mass and lean dry mass of skeletal muscles in black-capped chickadees: (a) body mass, (b) total lean dry muscle mass (including bones), (c) lean dry pectoralis muscle mass (filled circles: 27 °C, open circles: $-10$ °C).
or limits thermogenic capacity\textsuperscript{25}, its effect might result from system performance rather than from organ size. Mathieu-Costello \textit{et al.} could not be confirmed. Mathieu-Costello \textit{et al.} observed that muscovy ducklings (\textit{Cairina moschata}) were able to increase heat production before the onset of leg muscles shivering at temperatures below the lower critical temperature (but note that this may not be the case in black-capped chickadees\textsuperscript{74}). This was associated with an upregulation of avian uncoupling proteins (avUCP) in these same muscles, although the thermogenic role of avUCP (see text for details).

If larger muscles are not an absolute prerequisite for improving thermogenic capacity, then why are chickadees typically found with larger flight muscles in winter compared to summer [e.g. refs \textsuperscript{26,31}]? One possibility is that winter locomotion for active foraging and daily fattening during cold, short working days requires a different flight pattern leading to larger flight muscles. Given that muscle mass correlates positively with M\textsubscript{sum} at all temperatures, this hypothesis could also potentially explain why a number of individuals in our wild source population were found to maintain M\textsubscript{sum} levels above that required to guarantee intra-winter survival\textsuperscript{75} if these individuals were also the most active in that population.

In sum, our experimental data showed a clear influence of muscle mass on both maintenance energy costs, measured as BMR, and maximal thermogenic capacity, measured as M\textsubscript{sum}, in black-capped chickadees. Our results also showed that, although large muscles may be beneficial in terms of heat production capacity, an increase in muscle size is clearly not required to elevate M\textsubscript{sum} in these birds. Instead, improvement in thermogenic capacity appear to be related to cellular-level adjustments during cold-acclimation. The mechanisms underlying these adjustments deserves further study.

**Data Availability**

The datasets generated and analysed during the current study are available from the corresponding author upon request.

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| Organ | Treatment | F (df) | P | Adjusted R\textsuperscript{2} | BIC | ΔBIC |
|-------|-----------|--------|---|-----------------|-----|-----|
| Total muscles\textsuperscript{a} | 14.3 (1,25) | <0.001 | 31.7 (1,25) | <0.0001 | 0.33 | −26.85 | — |
| Pectoral | 12.0 (1,25) | <0.01 | 13.9 (1,25) | <0.001 | 0.32 | −25.2 | −1.65 |
| Legs\textsuperscript{a} | 10.7 (1,25) | <0.01 | 31.7 (1,25) | <0.0001 | 0.32 | −24.1 | −2.75 |
| Carcass\textsuperscript{a} | 10.3 (1,25) | <0.01 | 29.1 (1,25) | <0.0001 | 0.28 | −23.8 | −3.05 |
| Skin | 4.2 (1,25) | 0.05 | 20.5 (1,25) | <0.0001 | 0.24 | −18.5 | −8.35 |
| Brain | 3.8 (1,25) | 0.06 | 20.5 (1,25) | <0.0001 | 0.26 | −18.1 | −8.75 |
| Lungs | 2.9 (1,25) | 0.10 | 19.7 (1,25) | <0.001 | 0.19 | −17.2 | −9.65 |
| Pancreas | 1.7 (1,22) | 0.21 | 19.6 (1,22) | <0.001 | 0.19 | −15.9 | −10.95 |
| Gizzard | 0.92 (1,25) | 0.35 | 13.7 (1,25) | <0.01 | 0.24 | −15.2 | −11.65 |
| Kidneys | 0.1 (1,25) | 0.75 | 15.0 (1,25) | <0.001 | 0.17 | −14.3 | −12.55 |
| Heart (ventricle) | 1.0 (1,24) | 0.32 | 8.8 (1,24) | <0.01 | 0.15 | −13.5 | −13.35 |
| Intestine | 0.3 (1,24) | 0.61 | 12.0 (1,24) | <0.01 | 0.15 | −12.7 | −14.15 |
| Liver | 0.2 (1,17) | 0.65 | 15.1 (1,17) | <0.01 | −0.04 | −11.1 | −15.75 |

Table 4. Correlations between M\textsubscript{sum} and body composition. Results are from final ANCOVA models, including lean dry mass of organ and treatment as variables. All interactions were non-significant. \textsuperscript{*}Includes bone mass (see text for details).
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Acknowledgements

We are grateful to the Corporation de la Forêt d’Enseignement et de Recherche de Macpès, who granted us access to the field facilities for capturing the birds used in this study. We thank Mikael Jaffré, Ayméric Bodin and Gael Lafenêtre for their help in capturing the birds as well as Véronique Desrosiers, Claire Bottini and Karine Dubois for their help with dissections, organ processing and data entry. We are also grateful to Lyette Régimbald for the maintenance of captive birds and for food intake measurement as well as to Jonathan Coudé for his help with laboratory material. Alain Caron provided precious statistical advice. This work was funded by a Team Research Project grant from the Fonds Québécois de la Recherche sur la Nature et les Technologies (FRQNT) and by Discovery grants from the Natural Sciences and Engineering Research Council (NSERC) of Canada to F. V. and P.U.B. This work also benefited from a Leader Opportunity Fund award from the Canada Foundation for Innovation (CFI) to F. V. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
Author Contributions
M.S.M. collected all the data, M.S.M. and F.V. performed analyses. F.V., P.U.B. and M.S.M. designed the study. M.S.M. and F.V. wrote the paper.

Additional Information
Competing Interests: The authors declare no competing interests.

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