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**Abstract** Semi-jacketed lead-cored or copper-based homogenous rifle bullets are commonly used for hunting big game. Ever since their introduction in the 1990's, copper-based bullets have not been widely accepted by hunters due to limited supply, higher expense, and the perception that they exhibit inferior killing efficiency and correspondingly higher wounding rates. Here, we present data showing that animal flight distances for roe deer, red deer, brown bear, and moose dispatched with lead- or copper-based hunting bullets did not significantly differ from an animal welfare standardized animal flight distance based on body mass. Lead-cored bullets typical fragment on impact; in comparison, copper-based bullets retain more mass and expand more than their leaden counterparts. Our data demonstrate that the relative killing efficiency of lead and copper bullets is similar in terms of animal flight distance after fatal shots. Hunters that traditionally use lead bullets should consider switching to copper bullets to enhance human and environmental health.

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**Keywords (separated by '-')** Animal flight distance - Animal welfare - Hunting bullet expansion - Killing efficiency - Lead and copper ammunition - Wound ballistics

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**Footnote Information** **Electronic supplementary material** The online version of this article (<https://doi.org/10.1007/s13280-019-01171-4>) contains supplementary material, which is available to authorized users.

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2 **Unleaded hunting: are copper bullets and lead-based bullets**  
3 **equally effective for killing big game?**

4 Sigbjørn Stokke , Jon M. Arnemo, Scott Brainerd

5 Received: 10 December 2018/Revised: 14 February 2019/Accepted: 8 March 2019

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29 **INTRODUCTION**

30 Rifles using modern ammunition are used worldwide to  
31 cull or harvest wild mammals in order to manage popula-  
32 tions and provide recreational, commercial, and subsis-  
33 tence hunting opportunities. Lead (Pb) has been the metal


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of choice for making rifle projectiles since the earliest 34  
muzzleloaders were used for hunting. The reason is obvi- 35  
ous—lead is widely available, easily extracted from ore, 36  
simple to purify, and cheap to manufacture when compared 37  
to most other non-ferrous metals. It has a notably higher 38  
density (11.3 g/cm<sup>3</sup>) and much lower tensile strength 39  
compared to other metals available for manufacturing 40  
bullets. It is highly ductile, which allows for rapid expan- 41  
sion after impact to create large wound channels and is thus 42  
well-suited as a material for hunting projectiles (Almar- 43  
Næss 1985; MacPherson 1994; Guruswamy 2000). 44

As a non-toxic alternative to lead, rifle projectiles made 45  
of copper (Cu) and copper-zinc (Zn) alloys (tombac and 46  
brass) have been available since the 1990's. Copper is an 47  
essential element required to maintain homeostasis in 48  
vertebrates, even though too high or too low dietary intake 49  
can induce adverse health effects (Stern 2010). Copper is 50  
more expensive than lead but is less dense (8.96 g/cm<sup>3</sup>), 51  
although it is denser than most forms of steel (< 8.05 g/ 52  
cm<sup>3</sup>). Lead is about 1.5 times more ductile than copper 53  
(Almar-Næss 1985). 54

**Bullet expansion and wound ballistics** 55

Hunting bullets designed to expand or deform will exhibit a 56  
mushroom-like anterior enlargement of the cross-sectional 57  
area of the bullet at impact. Lead-based hunting bullets (L- 58  
bullets) have a lead core covered with a copper jacket 59  
except for the leading lead tip. At impact the lead core 60  
behaves like an incompressible fluid when the drag forces 61  
generated by the stagnation pressure at the leading edge of 62  
the bullet exceed the yield limit for lead (Berlin et al. 1988; 63  
MacPherson 1994; Kneubuehl et al. 2011). Pressure is thus 64  
dispersed within the floating lead and works the jacket 65  
from the inside of the bullet, causing it to burst (Berlin 66

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67 et al. 1988; Kneubuehl et al. 2011). Expansion is very rapid  
68 and stagnates within 0.1 ms (Kneubuehl et al. 2011).  
69 Copper-based, homogeneous lead-free hunting bullets (C-  
70 bullets) expand according to the same mechanisms if the  
71 frontal cavity is large enough for viscous pressure to enter  
72 (Kneubuehl et al. 2011).

73 Bullet penetration is characterized by the temporary  
74 cavity caused by tissue impelled radially in relation to the  
75 velocity vector as momentum is imparted from the pro-  
76 jectile to the soft tissue and it undergoes elastic deformation  
77 as it is stretched and compressed (Stefanopoulos et al.  
78 2014). The displaced tissue will rapidly recoil towards its  
79 initial position in response to the vacuum and elastic  
80 energy conveyed to the tissue, thus generating a brief  
81 oscillation (Harvey et al. 1946; Di Maio 1999; Fackler  
82 2001; Kneubuehl et al. 2011). The residual wound channel,  
83 which is a cavity filled with blood, damaged tissue, and  
84 contaminants sucked in from the outside, is termed the  
85 permanent wound cavity (Fackler 1988; Janzon et al.  
86 1997). The extravasation zone is the transition between the  
87 permanent wound cavity and intact tissue and is charac-  
88 terized by hemorrhage resulting from distention of the  
89 temporary cavity, inflicting damage to blood vessels through  
90 overstretching and shearing effects due to heterogeneity of  
91 the involved tissues (Kneubuehl et al. 2011; Stefanopoulos  
92 et al. 2014). There is a proportional relationship between  
93 the kinetic energy of the penetrating bullet and the  
94 expansion of the temporary cavity. Thus, the potential  
95 energy stored in the tissue equals the work done to create  
96 the maximum expansion. MacPherson (1994) states that  
97 the potential for this energy to cause wounding depends on  
98 four factors: The magnitude of the stored energy in the  
99 tissue, the ability of the tissue to sustain strain, the size of  
100 the organ structure, and the anatomical constraints to tissue  
101 movements. If the energy stored in tissue exceeds the  
102 elastic limit of the tissue, it will rupture and permanent  
103 wounding results. Tissue elasticity is therefore an impor-  
104 tant factor as it impairs the extent of permanent damage  
105 caused by a bullet. Muscle, skin, blood vessels, and lungs  
106 are elastic and can absorb energy generated by a pen-  
107 etrating bullet and tend to recoil towards the wound channel  
108 (Fackler 1988; MacPherson 1994; Karger 2008). Other less  
109 resilient tissues, such as liver, kidney, and brain, tend to  
110 disrupt from penetrating projectiles (Roberts 1988; Caudell  
111 2013; Stefanopoulos et al. 2014).

112 The size of the organ or body is important because there  
113 will be a lower size limit whereby all tissues will be  
114 stretched beyond the elastic limit of the organ or body,  
115 causing it to rupture. For organs or bodies larger than this  
116 critical size, tissue damage primarily occurs by crushing,  
117 tearing, and stress (MacPherson 1994). Thus, the primary  
118 factor causing permanent wound cavity in soft tissue like  
119 lungs will mainly be crushing rather than radial stretching

if the organ size exceeds the critical size (Stefanopoulos 120  
et al. 2014). This suggests that the area of the leading edge 121  
of the bullet might correlate with the radial dimension of 122  
the permanent wound cavity, with larger calibers yielding 123  
larger wound channels. Fragmentation is an inherent ability 124  
of all lead-based bullets where lead floats and expands in 125  
response to the stagnation pressure (Fackler et al. 1984; 126  
Cornicelli and Grund 2008; Stokke et al. 2017). Although 127  
debated, bullet fragmentation is commonly considered to 128  
be a primary cause of increasing the permanent wound 129  
cavity by weakening the tissues under tension from the 130  
temporary cavity (Fackler et al. 1984; Coupland 1999; 131  
Trinogga et al. 2013). In contrast, deforming copper bullets 132  
can withstand fragmentation and thus sustain momentum 133  
ensuring proper penetration (Hunt et al. 2009; Batha and 134  
Lehman 2010; Gremse et al. 2014). 135

### Cause of death for animals dispatched with hunting bullets 136 137

Most hunters, in accordance to codes of practice, target the 138  
thoracic area. The expanded bullet will penetrate the tho- 139  
racic cavity, causing trauma to the heart, lungs, and/or 140  
major blood vessels causing subsequent fatal hemorrhage 141  
(with subsequent hypotension, hypovolemic shock, 142  
hypoxia, and brain death) (Stokke et al. 2018). Hemorrhage 143  
is the cause of death in hunted animals, unless the bullet 144  
traumatizes the brain (brain death) or the spinal cord cran- 145  
ial to C3–C5 (where the phrenic nerves exit). Wounded, 146  
immobile animals are dispatched (euthanized) with a 147  
head/neck shot and then the cause of death is not fatal 148  
bleeding. Impacts to other body parts might cause fatal 149  
hemorrhaging if large blood vessels are lacerated or a well- 150  
perfused organ such as a kidney or the liver is ruptured. 151  
Fatal wounds will inevitably be followed by circulatory 152  
collapse due to a hypovolemic shock with subsequent brain 153  
hypoxia (Vincent and De Backer 2013; Gaieski and Mik- 154  
kelsen 2017). Death due to blood loss is never instantane- 155  
ous and the rate of hemorrhaging determines the time 156  
from bullet impact to permanent incapacitation. Therefore, 157  
animal flight distance conveys information about elapsed 158  
time and can be used as a practical indicator for killing 159  
efficiency of hunting bullets and cartridges (Stokke et al. 160  
2012; McCann et al. 2016; Kanstrup et al. 2016b; Martin 161  
et al. 2017; Stokke et al. 2018). 162

### Lead toxicity and transition to non-lead ammunition 163

Even though the use of L-bullets is mainstream, there are 164  
concerns over health and environmental risks from spent 165  
ammunition (Bellinger et al. 2013). Lead has no known 166  
biological function in vertebrates and is toxic to most 167  
physiological systems (Bellinger et al. 2013). A transition 168

169 to C-bullets is therefore strongly recommended to avoid  
 170 lead exposure in humans consuming game meat and in wild  
 171 animals scavenging on remains from shot game (Krone and  
 172 Hofer 2005; Grund et al. 2010; Delahay and Spray 2015;  
 173 Arnemo et al. 2016; Kanstrup et al. 2016a; McTee et al.  
 174 2017; Gerofke et al. 2018; Kanstrup et al. 2018). In contrast  
 175 to lead, copper is an essential element in vertebrates and is  
 176 generally not considered to be toxic to humans (Stern  
 177 2010).

178 Hunters have raised concerns over the efficiency of  
 179 C-bullets (Caudell et al. 2012; Bundesinstitut für  
 180 Risikobewertung 2013), including the perception of limited  
 181 supply, higher costs, inferior killing efficiency, and corre-  
 182 spondingly higher wounding rates compared to 'traditional'  
 183 lead-based ammunition (Southwick Associates Inc. 2014;  
 184 Thomas et al. 2016). However, C-bullets compare favor-  
 185 ably to L-bullets in recent studies. In a controlled experi-  
 186 ment, Gremse et al. (2014) used ballistic soap as tissue  
 187 simulant to show that the terminal ballistics of C- bullets  
 188 were similar to L-bullets. However, tissue simulants are  
 189 very different from live tissue and may not be analogous to  
 190 living animals. Trinogga et al. (2013) examined 34 car-  
 191 casses of ungulates [wild boar (*Sus scrofa*), roe deer  
 192 (*Capreolus capreolus*), chamois (*Rupicapra rupicapra*),  
 193 and red deer (*Cervus elaphus*)] shot with either L- or  
 194 C-bullets. They used X-ray computed tomography to  
 195 measure permanent wound cavities in the lungs and con-  
 196 cluded that both bullet types should have the same killing  
 197 potential. However, if hunters are to use C-bullets with  
 198 confidence, they want to see evidence from actual hunting  
 199 situations where uncontrolled events may occur. Kanstrup  
 200 et al. (2016b) conducted a study that included 657 ungu-  
 201 lates shot with either L- bullets or C-bullets by recreational  
 202 hunters. The authors used animal flight distance as the  
 203 primary response variable and concluded that C-bullets  
 204 were an effective alternative to L-bullets. Spicher (2008)  
 205 found that 95% of 247 animals were killed quickly with a  
 206 single shot from C-bullets. Of the 12 hunters in that survey,  
 207 eight (66%) were convinced that C-bullets were as suit-  
 208 able as traditional L-bullets, and four (33%) considered that  
 209 the C-bullets performed better. Knott et al. (2009) studied  
 210 red deer and roe deer dispatched with either C- or L-bul-  
 211 lets. They reported no significant difference between either  
 212 bullet type regarding killing efficiency or accuracy. Like-  
 213 wise, McCann et al. (2016) found that C-bullets were  
 214 effective in culling 983 elk (*Cervus elaphus*). Finally,  
 215 McTee et al. (2017) studied the capacity of L- and  
 216 C-bullets to instantly incapacitate ground squirrels (*Sci-*  
 217 *uridae* spp) and found no difference between the two bullet  
 218 types.

## Hypotheses and objectives

220 We used animal flight distance as a discriminator to study  
 221 differences in killing efficiency between expanding L- and  
 222 C-bullets. In doing so, we applied the new model devel-  
 223 oped by Stokke et al. (2018) to compare observed animal  
 224 flight distances with animal flight distance welfare stan-  
 225 dards for Fennoscandia (Stokke et al. 2018). This model  
 226 estimates an expected animal flight distance for mammals  
 227 based on body mass. One advantage of using this model is  
 228 its objective representation of animal welfare outcomes  
 229 that reflect physiological processes that occur in an animal  
 230 during and after bullet penetration. Furthermore, the model  
 231 enables a comparison of animal flight distances without  
 232 dividing the data into groups based on mammal species,  
 233 body mass, or age classes. In addition, we developed  
 234 indices for bullet expansion and degree of asymmetrical  
 235 expansion to study differences in expansion potential  
 236 between the two bullet types. We tested the null hypothesis  
 237 that both bullet types exhibited similar killing efficiency  
 238 and expansion characteristics.

## MATERIALS AND METHODS

### Sampling of hunting data

241 Big game hunting in Fennoscandia is typically performed  
 242 in hunting teams including around 6 hunters on average in  
 243 Norway (Solberg et al. 2014). During hunting, team  
 244 members position themselves at strategic sites where ani-  
 245 mals predictably pass by when driven by other hunters with  
 246 or without the aid of hunting dogs. In these circumstances,  
 247 shooting distances are usually within 100–150 m (Fig. 1).

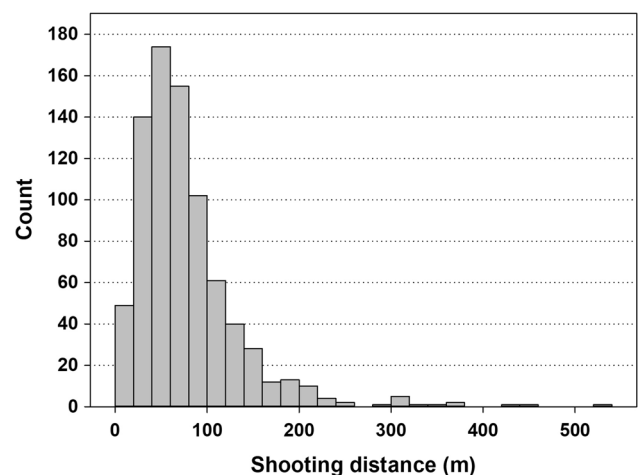


Fig. 1 Frequencies of shooting distances recorded during the present study

248 We collected data from four mammalian species based  
 249 on questionnaires distributed to hunters in Fennoscandia:  
 250 Moose (*Alces alces*: Finland, Sweden, and Norway  
 251 2004–2006)  $n = 5\ 245$ ; brown bear (*Ursus arctos*: Sweden  
 252 2006–2010)  $n = 637$ ; roe deer (Norway 2014–2015)  
 253  $n = 38$ ; red deer (Norway 2014–2015)  $n = 1$ . The hunters  
 254 completed one form per harvested animal. Live body  
 255 masses ranged from 9 kg (roe deer) to 662 kg (moose). In  
 256 addition, hunters provided bullets retrieved from moose  
 257 carcasses together with the corresponding questionnaire  
 258 ( $n = 1833$ , see Online Appendix S1 for summary of bullet  
 259 types).

260 In this paper, we used the following information from  
 261 the questionnaires: animal flight distance (m), number of  
 262 impacting bullets, whether the bullet exited or stopped in  
 263 the animal body, the angle of the bullet trajectory in rela-  
 264 tion to the animal's longitudinal axis, penetrated organs  
 265 and bones, cartridge, bullet type, whole or slaughter mass,  
 266 and age class (moose only). We discerned between  
 267 L-bullets and C-bullets as defined above. To avoid skew-  
 268 ness in animal flight distances due to caliber size, we  
 269 included only calibers with both L- and C-bullets in the  
 270 analyses. The most commonly used calibers ranged from  
 271 6.5 to 8.0 mm.

272 For all roe deer, red deer, moose, and some bears, we  
 273 converted slaughter weights ( $W_s$ ) to estimated live masses  
 274 ( $M_1$ ) (kg). For roe deer, red deer, and moose, we estimated  
 275 this using the following formula (Hjorteviltregisteret  
 276 2016):

$$M_1 = \frac{100 \cdot W_s}{52}$$

278 For bears, we estimated live masse ( $M_b$ ) (Swenson et al.  
 279 1995) using the formula

$$M_b = 4.63 + 1.49 \cdot W_s$$

281 Hunters were asked to locate the spot where the animal  
 282 was struck by the first bullet and from that point start  
 283 pacing out along the track of the animal until they arrived  
 284 at the incapacitated animal. This route, covered by the shot  
 285 animal, was recorded as animal flight distance in the form.

## 286 Comparison of efficiency of lead-based 287 versus homogenous bullets

288 Concerns have been raised regarding the performance of  
 289 bullets, in particular C-bullets, when shooting distance  
 290 exceeds 200 m (Caudell et al. 2012; Caudell 2013). Caudell  
 291 et al. (2012), firstly draws attention to the possibility of  
 292 destabilized bullets due to a mismatch between bullet  
 293 length and twist rate of the rifle barrel, and secondly to  
 294 reduced expansion potential. Even though shooting dis-  
 295 tances in the present study rarely exceeded 150 m, we

296 examined if expansion was affected within recorded  
 297 shooting distances to ensure that our modeling was not  
 298 influenced by this factor. Due to very few records for  
 299 shooting distances exceeding 150 m (Fig. 1), we excluded  
 300 records for longer shooting distances. We applied our  
 301 expansion indices for this purpose (see next chapter). We  
 302 regressed the indices against shooting distances and  
 303 exhibited the result in scatter diagrams with a linear  
 304 regression per bullet and caliber category.

305 To enable a sound comparison between animal flight  
 306 distances shot with C- or L-bullets, we included only  
 307 records fulfilling the following requirements: (1) the target  
 308 animal was dispatched with one bullet; (2) the bullet tra-  
 309 jectory described an angle of incidence  $\leq 45^\circ$  (in relation  
 310 to the longitudinal axis of the animal in the horizontal  
 311 plane), (3) bullet type and caliber were known and, (4) both  
 312 lungs were penetrated. These criteria reduced the number  
 313 of records to 710 moose, 71 bears, 1 red deer, and 32 roe  
 314 deer.

315 To evaluate if any discrepancies existed between animal  
 316 flight distances caused by C- or L-bullets, we applied the  
 317 model designed by Stokke et al. (2018) defining animal  
 318 welfare standards in hunting (Fig. 2). Based on penetration  
 319 of the thoracic region, the model estimates an expected

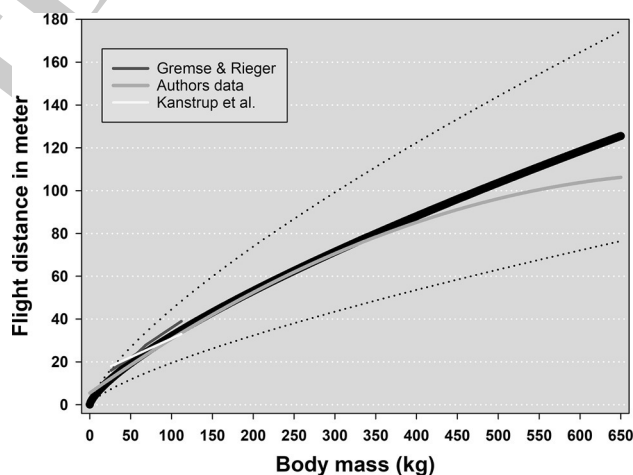


Fig. 2 Expected animal flight distances (efd) predicted by the model for mammals with body masses < 650 kg (reprinted from Stokke et al. 2018). The solid broad black line represents efd in relation to body mass. Dotted lines represent the uncertainty of parameter estimation. A very good accordance with average animal flight distances, recorded from several mammal species under field hunting conditions, exhibits the predictive power of the model. The dark short grey line displays average animal flight distances for four species with increasing body masses: roe deer, fallow deer (*Dama dama*), wild boar (*Sus scrofa*), and red deer (Gremse and Rieger 2014). The short white line shows average animal flight distances for roe and red deer (Kanstrup et al. 2016b). The long light grey line is the regression line representing animal flight distances for red fox, roe deer, brown bear, and moose calves, yearlings, and adults (Stokke et al. 2018)

320 animal flight distance (efd), for mammals if body mass  
321 ( $M$ ) is known (Stokke et al. 2018).

$$\text{efd} = 1.14M^{0.73}$$

323 In the applied form the model is calibrated with  
324 estimated average traveling speed for adult moose after  
325 being shot. However, it is obvious that traveling speed for  
326 animals penetrated by expanding bullets may vary. To  
327 compensate for this, the model can be calibrated with  
328 estimated speed for the species in question. Here we apply  
329 four mammalian species. So, the question is, did the  
330 animals travel with sufficiently equal speed (i.e., similar  
331 deviations from estimated efd-values) to justify a  
332 comparison without addressing differences among  
333 species? In our case this could partly be tested, because  
334 brown bear and moose (calves, yearlings, and adults) had  
335 enough overlap of body masses to address this question.  
336 We applied records for brown bear and all records from  
337 moose except those representing body masses outside the  
338 range of brown bear body masses. The data were grouped  
339 into 5 body mass classes representing a stepwise increase  
340 of 50 kg per class (range 38–250 kg). We exhibited  
341 deviations from efd-values with error bars and used a  
342 general linear model to test for speed differences between  
343 the species.

344 For all body masses and species, we calculated the  
345 discrepancies between expected animal flight distances  
346 (efd) and reported animal flight distances and conveyed the  
347 differences into two samples (1 and 2) according to bullet  
348 class [L ( $n_1 = 729$ ) vs C ( $n_2 = 84$ )]. These samples were  
349 compared using a general bootstrap approach with ran-  
350 domized residuals (Ter Braak 1992; Manly 2001). First, we  
351 computed the  $t$ -statistics for these samples. This  $t$ -value  
352 was then compared with a bootstrap distribution for which  
353 the null hypothesis was made to be true by replacing the  
354 sample values with their residuals. A bootstrap population  
355 of residuals of size  $n_1 + n_2$  could then be used to draw a  
356 bootstrap sample 1 by selecting  $n_1$  of these values at ran-  
357 dom with replacement. Similarly, we obtained a bootstrap  
358 sample 2 by selecting  $n_2$  cases from the bootstrap popu-  
359 lation. These samples were used to compute a bootstrap  
360 value for  $t$ . By repeating this procedure many times, the  
361 bootstrap distribution of  $t$  was generated. A two-sided test  
362 was applied to see if the  $|t|$  value for the observed samples  
363 (1 and 2) was significantly larger compared to the distri-  
364 bution of bootstrapped  $|t|$  values from randomized residu-  
365 als. Accordingly, this test does not produce any  $p$  value.

366 To examine if discrepancies between estimated and  
367 recorded animal flight distances related to C- or L-bullets  
368 differed among body masses, we pooled body masses into  
369 weight classes divided per 50 kg body mass up to 200 kg.  
370 Due to few samples for C-bullets related to body  
371 masses  $> 200$  kg, we applied two weight classes between

200 and 650 kg. The result was exhibited in a grouped  
vertical error bar graph and tested with a general linear  
model.

Furthermore, we applied the animal welfare standard  
model to compare the data against the wounding threshold  
or maximal animal flight distance (mfd) suggested by the  
model (Stokke et al. 2018).

$$\text{mfd} = 4.92M^{0.73}$$

### Bullet expansion index and penetration ability

Retrieved bullets were processed according to Stokke et al.  
(2017). The expanded frontal area of the bullets was  
measured using a Vernier caliper to obtain two cross-sec-  
tional measurements ( $d_1$  and  $d_2$ —perpendicularly oriented  
to each other) to even out asymmetrical expansion. These  
measurements were used together with bullet diameter ( $d$ )  
to express an index ( $E$ ) for rate of expansion in relation to  
original bullet cross-sectional area.

$$E = \left( \frac{d_1 + d_2}{2d} \right)^2$$

This expansion index was used to explore the expansion  
potential of C- versus L-bullets and to analyze if caliber  
and expansion index are correlated. We divided bullets (L  
and C) into three caliber categories according to diameter  
(mm): (1) 6.5–7.9, (2) 8.0–9.8, and (3)  $> 9.9$ . Variation of  
expansion among these categories for L- and C- bullets was  
exhibited with an error bar graph and tested with a general  
linear model. Due to very low sample size for C-bullets in  
the largest caliber category, this category was excluded  
from the statistical model.

In addition, we express asymmetrical expansion with the  
following index:

$$E_{\text{sym}} = 1 - \left( \frac{d_s}{d_l} \right)^{-1}$$

where  $d_s$  represents the smallest and  $d_l$  the largest of the  
two diameters  $d_1$  and  $d_2$ . This index equals zero if the  
diameters are alike and decreases linearly with increasing  
differences between the diameters. We used this index to  
compare the levels of asymmetrical expansion between C-  
and L-bullets and tested for a difference with the Mann-  
Whitney  $U$  test.

Finally, we studied the ability of bullets to fully pene-  
trate and exit moose bodies. This was done by calculating  
the ratio between bullets that exited the bodies and those  
that stopped in the bodies. We divided the analyses into  
two caliber categories: 6.5–7.9 mm and  $\geq 8.0$  mm. Due to  
reasonable sample sizes for smaller calibers, we were able  
to divide L-bullets into three categories: bonded core  
(copper jacketed soldered to the lead core), h-mantel (dual



419 lead cores separated with integral partitioning of the copper  
 420 jacket), and conventional (simple copper jacket with  
 421 unsoldered lead core). Furthermore, we divided moose  
 422 body sizes into three age categories for this analysis:  
 423 calves, yearlings, and adults. We applied a generalized  
 424 linear model with binomial distribution to analyze pen-  
 425etration ability. This was done only for the smallest caliber  
 426 category due to very few samples for the larger calibers.

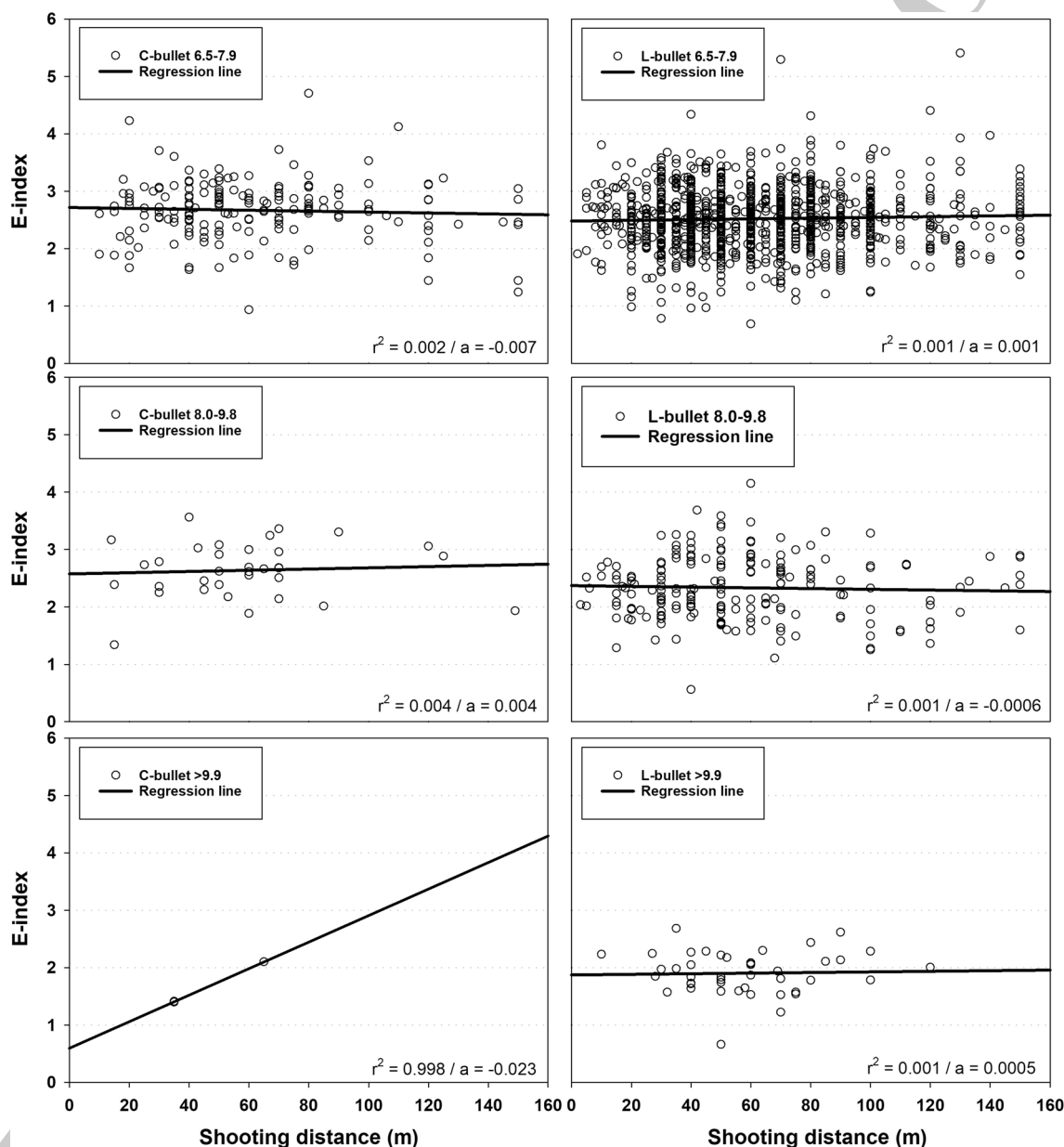
427 We used Visual FoxPro 9.0 SP2 to handle the data and  
 428 to program the bootstrap session. We performed standard

statistical analyses with IBM SPSS Statistics Version 25  
 and created graphs with SigmaPlot 13.0.

**RESULTS**

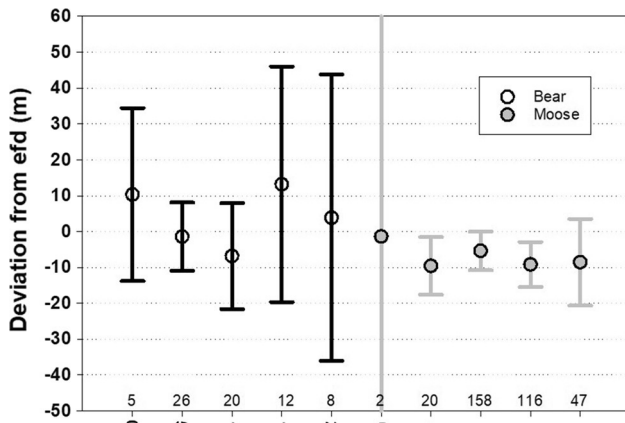
**Comparison of killing efficiency for C- and L-bullets**

Apparently, expansion rate of bullets was unaffected by  
 shooting distance within 150 m, except for C-bullets in  
 the > 9.9 mm caliber category (Fig. 3). However, nothing



**Fig. 3** Expansion indices for C- and L-bullets within three caliber categories in relation to shooting distance, exhibited from top to bottom: (1) 6.5–7.9 mm, (2) 8.0–9.8 mm, and (3) > 9.9 mm. C-bullets are on the left and L-bullets to the right. Values for  $r^2$  and slope (a) are depicted at the bottom of each graph

Author Proof



**Fig. 4** Deviations from predicted edf-values for recorded animal flight distances from brown bear and moose exhibited in 5 body mass classes (kg). The number of records per body mass class and species is displayed below corresponding error bars

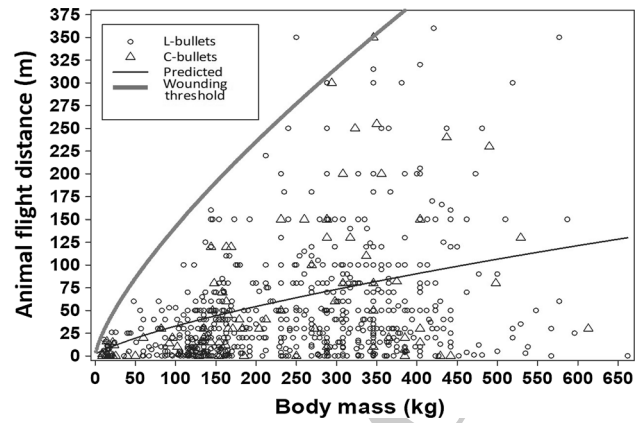
436 can be deduced from this regression due to lack of data-  
 437 points for this category. Since bullets exhibited constant  
 438 expansion potential within shooting distances shorter than  
 439 150 m, we did not regard this factor to have any significant  
 440 effect on our approach to examine killing efficiency.

441 Recorded animal flight distances for brown bear and  
 442 moose exhibited similar deviations from predicted edf-  
 443 values among body mass groups (Fig. 4:  $F = 0.40$ ,  $df = 4$ ,  
 444  $p = 0.81$ ). Furthermore, deviations did not differ between  
 445 brown bear and moose (Fig. 4:  $F = 2.12$ ,  $df = 1$ ,  $p = 0.15$ ).  
 446 There was no interaction between deviations for brown  
 447 bear and moose (Fig. 4:  $F = 0.65$ ,  $df = 4$ ,  $df_{error} = 387$ ,  
 448  $p = 0.63$ ). This suggests that traveling speed following  
 449 bullet impact for these species was analogous and unlikely  
 450 to skew model output and statistical analyses noticeably.

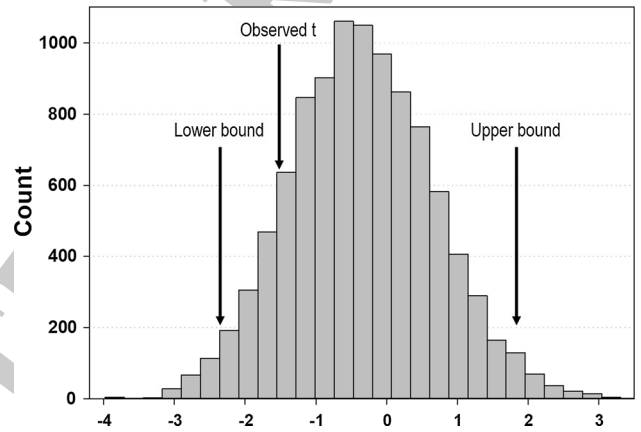
451 Measured animal flight distances exhibited a large  
 452 variation in relation to predicted animal flight distances  
 453 (Fig. 5). Yet, most records were reasonably evenly dis-  
 454 tributed around the expected animal flight distances  
 455 (Fig. 5). Recorded animal flight distances, with one  
 456 exception, were below the wounding threshold suggested  
 457 by the model (Stokke et al. 2018). Actual animal flight  
 458 distances exhibited increasing variability with body mass.

459 The bootstrap approach suggested that deviations from  
 460 the expected animal flight distances (efd) did not differ  
 461 between animals dispatched with L- or C-bullets (Fig. 6).  
 462 This is because the  $t$ -value for the observed data is located  
 463 within the confidence interval for the bootstrapped  $t$ -values  
 464 from randomized residuals (Fig. 6).

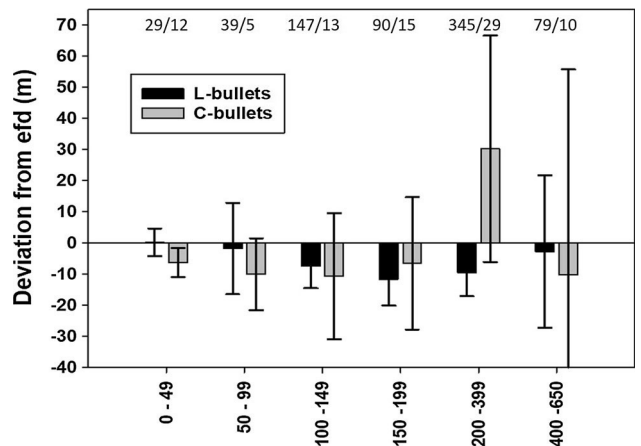
465 Recorded deviations from predicted animal flight dis-  
 466 tances did not vary significantly among body masses  
 467 (Fig. 7:  $F = 0.69$ ,  $df = 5$ ,  $df_{error} = 5$ ,  $p > 0.6$ ). It might be  
 468 worthwhile noting that C-bullets on average gave shorter



**Fig. 5** Distribution of animal flight distances of C- and L-bullets compared to predicted animal flight distances (thin lower line) and wounding threshold (bold upper line) derived from the model (Stokke et al. 2018)



**Fig. 6** Randomized residual bootstrap distribution of  $t$ -values compared to the  $t$ -value ( $-1.65$ ) for the observed data. The analysis is performed with 10 000 bootstrap samples with mean =  $-0.36$ , Lower bound =  $-2.33$  and Upper bound =  $1.73$



**Fig. 7** Deviation from predicted animal flight distances, according to edf-values (Stokke et al. 2018), for C- and L-bullets in relation to 6 body mass classes. Sample sizes are exhibited above the vertical error bars (L-bullets/C-bullets)

**Table 1** Range and variance of expansion indices for C- and L-bullets within three caliber categories

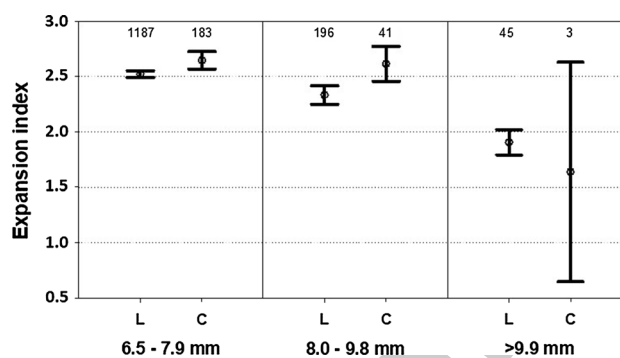
Bullet	Caliber category (mm)	N	Range	Variance
C	6.5–7.9	183	3.77	0.29
	8.0–9.8	41	3.56	0.40
	> 9.9	3	0.69	0.16
L	6.5–7.9	1187	5.40	0.47
	8.0–9.8	196	4.74	0.42
	> 9.9	45	2.68	0.29

469 animal flight distances for the three smallest body mass  
470 categories, whereas the situation was reversed for the next  
471 two categories. However, this shift of deviation from efd-  
472 values was not significant (Fig. 7:  $F = 0.10$ ,  $df = 1$ ,  $df_{er-}$   
473  $ror = 6.87$ ,  $p > 0.7$ ). There was no interaction between C-  
474 and L-bullets and body mass classes regarding deviation  
475 from predicted animal flight distance (Fig. 7:  $F = 1.43$ ,  
476  $df = 5$ ,  $df_{error} = 801$ ,  $p > 0.2$ ). However, due to small  
477 sample sizes for C-bullets, these results should be treated  
478 with caution.

#### 479 Comparison of expansion indices and penetration 480 ability

481 Within the three caliber categories, L-bullets appeared to  
482 have a wider range of expansion compared to C-bullets  
483 (Table 1). This indicates that L-bullets exhibited a more  
484 irregular expansion history than C-bullets. This is also  
485 reflected in the variance, which is smaller for C-bullets,  
486 irrespective of smaller sample sizes (Table 1.). This  
487 expansion pattern is supported by the Levene's test, sug-  
488 gesting unequal variances (Levene statistics: mean = 2.52,  
489  $df_1 = 5$ ,  $df_2 = 1604$ ,  $p = 0.02$ ).

490 The expansion potential was apparently largest for  
491 C-bullets as their indices were larger than indices of  
492 L-bullets (Fig. 8:  $F = 20.3$ ,  $df = 1$ ,  $df_{error} = 1604$ ,  
493  $p < 0.001$  {> 9.9 mm category not included}). In the  
494 smallest caliber category, C-bullets expansion index was  
495 on average 2.65 compared to 2.52 for L-bullets (Fig. 8).  
496 The same trend was evident for the next caliber category  
497 with an index of 2.62 for C-bullets versus 2.33 for L-bullets  
498 (Fig. 8). For the largest caliber category, the trend was  
499 reversed, and C-bullets exhibited an index of 1.64 versus  
500 1.91 for L-bullets. However, sample size for C-bullets in  
501 the last category is very small and thus the comparison is  
502 unreliable. Another expansion trend was the capacity of  
503 C-bullets to maintain expansion index when caliber size  
504 increased from the smallest caliber category to the medium  
505 one (Fig. 8). This trend was absent for L-bullets as they  
506 exhibited a steady decrease of expansion indices for  
507 increasing caliber (Fig. 8).



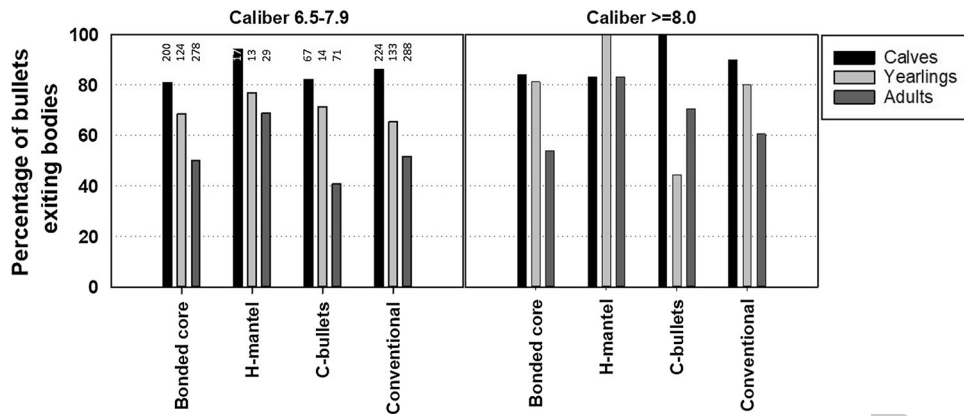
**Fig. 8** Expansion indices for C- and L-bullets within three caliber categories. Sample sizes are shown above their respective error bars

The index ( $E_{sym}$ ) representing the level of asymmetrical  
expansion showed that L-bullets expanded more asym-  
metrically than C-bullets (Mann–Whitney  $U = 136\ 516.5$ ,  
 $p = 0.002$ ). Average asymmetrical index for C-bullets was  
 $-0.09$ , whereas corresponding index for L-bullets was  
 $-1.13$ .

The tendency of bullets to exit moose bodies did not  
vary among bullet categories in the smallest caliber cate-  
gory (Fig. 9: Wald Chi-Square = 4.74,  $df = 3$ ,  $p = 0.2$ ). All  
bullet types exhibited a clear tendency to increase the  
amount of seizures with increasing body size (Fig. 9: Wald  
Chi Square = 46.83,  $df = 2$ ,  $p < 0.001$ ). This pattern was  
consistent for all bullet categories in the smallest caliber  
category as no interaction was present between age classes  
and bullet categories (Fig. 9: Wald Chi Square = 3.36,  
 $df = 6$ ,  $p = 0.8$ ).

## DISCUSSION

An evaluation of the efficacy of non-lead versus lead-based  
ammunition has never been done based on quantified ani-  
mal welfare outcomes. In this paper, we applied a novel  
model designed by Stokke et al. (2018) that defines humane  
killing. The model predicts animal flight distances fol-  
lowing penetration of both lungs in relation to body mass  
of mammals. From an animal welfare perspective, the  
targeting of vital organs is the optimal and most humane  
killing strategy because it induces rapid and fatal hemor-  
rhaging (Stokke et al. 2018). In our approach, we measured  
deviations for animal flight distances recorded by hunters  
with standardized animal flight distances (efd) suggested  
by the model in relation to body mass. In contrast to other  
studies, our approach allows a direct comparison of animal  
flight distances without any need to classify animals into  
groups according to body size, age class, or species. This is  
because the model is mathematically deduced from allo-  
metric relationships generally acknowledged to be



**Fig. 9** The percentage of bullets exiting moose bodies in relation to age class, bullet, and caliber category. Bonded core, H-mantel, and Conventional belong to L-bullets. Sample sizes are displayed above their respective bars. Sample sizes for the largest caliber category are too small for statistical analyses

543 universal for mammals. However, if studied mammalian  
 544 game species travel with unequal speeds after bullet  
 545 impact, the result might be skewed, and corrections should  
 546 be applied (Stokke et al. 2018). Brown bear and moose had  
 547 the largest overlap of body masses so we could test for  
 548 differences between traveling speed for these species. We  
 549 did not detect significant speed differences (i.e., equal  
 550 deviations from efd-values) between brown bear and  
 551 moose, thus indicating no need to differ among species  
 552 during the analyses.

553 Our findings showed that animal flight distances varied  
 554 greatly although we only used cases where bullets pene-  
 555 trated both lungs. Variability also increased with body  
 556 mass. This pattern is to be expected because (1) total blood  
 557 volume remains unchanged in relation to body mass, (2)  
 558 blood circulation time increases with body mass, whereas  
 559 (3) the radial dimension of the permanent wound cavity  
 560 remains largely unchanged (Stokke et al. 2018). Thus,  
 561 bleeding rates will decrease, whereas animal flight distances  
 562 will increase with increasing body mass. The model esti-  
 563 mates animal flight distances relative to body mass when an  
 564 animal is dispatched with an expanding bullet penetrating  
 565 both lungs centrally and perpendicularly to its longitudinal  
 566 axis. Thus, peripheral penetrations of the thorax area will  
 567 yield diminished hemorrhaging followed by increased ani-  
 568 mal flight distances because less lung tissue is disrupted.  
 569 We believe that the model adequately estimates optimal  
 570 exsanguination rates in relation to body mass. However,  
 571 target animals of equal body mass after bullet impact may  
 572 travel at different velocities, which will affect animal flight  
 573 distances and increase variability of animal flight distances.  
 574 Interestingly, it appears that the wounding threshold (mfd)  
 575 defined by the model clearly delineates all cases, except  
 576 one, from the region defining wounding. This suggests that  
 577 the model is appropriate to evaluate killing of animals in  
 578 relation to animal welfare standards.

579 There was no significant difference in animal flight  
 580 distances among animals (moose, brown bear, roe deer, and  
 581 red deer) incapacitated with L- or C-bullets when com-  
 582 pared with efd-values in relation to body mass. For all body  
 583 mass classes, deviations from predicted efd-values for C-  
 584 and L-bullets were below and close to efd-values. How-  
 585 ever, for the 200–399 kg body mass class, deviations from  
 586 predicted efd-values for C-bullets were above predicted  
 587 efd-values, whereas corresponding deviations for L-bullets  
 588 were below predicted ones. The difference between the two  
 589 bullet types was insignificant, but never the less noticeable  
 590 and might be of interest for hunters, but sample size for  
 591 C-bullets was low and the discrepancy might as well be  
 592 coincidental.

593 Kanstrup et al. (2016b) noticed a similar tendency for  
 594 animal flight distances recorded from dispatched roe deer  
 595 when shooting distances exceeded 100 m. There is one  
 596 obvious difference in expansion history for the two bullet  
 597 types. L-bullets retrieved from moose carcasses lose on  
 598 average 2.8 g of lead per bullet, whereas C-bullets loose  
 599 around 0.5 g of mass (copper) per bullet (Stokke et al. 2017).  
 600 Fragmentation is therefore much more pronounced for  
 601 L-bullets. These fragments might enlarge the bleeding sur-  
 602 face of the wound cavity by penetrating and weakening  
 603 tissue in the extravasation zone during cavitation and thus  
 604 enhance rupturing of tissue (Fackler et al. 1984). The  
 605 importance of enhanced hemorrhage in the extravasation  
 606 zone is also noted by Stokke et al. (2018), as they suggested  
 607 that this zone is a functional part of the wound. So, how can  
 608 non-fragmenting C-bullets compete so well with L-bullets  
 609 that have this inherent advantage of fragmentation?

610 Our results suggest two areas where C-bullets outper-  
 611 formed L-bullets. Firstly, they expanded more and pre-  
 612 sented a larger frontal surface after tissue penetration than  
 613 L-bullets. For the most common caliber categories  
 614 (6.5–7.9 mm and 8.0–9.8 mm), C-bullets exhibited a

615 stable expansion index around 2.6, whereas L-bullets barely  
 616 reached 2.5 within the smallest caliber category  
 617 (6.5–7.9 mm) and diminished strongly for larger caliber  
 618 categories. Secondly, C-bullets exhibited a more consistent  
 619 and stable expansion than L-bullets. Both range and variance  
 620 were consistently less for expansion indices within all caliber  
 621 categories for C-bullets compared to L-bullets. This clean-  
 622 cut expansion pattern for C-bullets compared to L-bullets is  
 623 probably related to their mechanical properties. Copper is  
 624 relatively ductile and deforms plastically when yielding.  
 625 However, C-bullets will not expand if not “weakened” by an  
 626 axial cylindrical hole in the anterior part of the bullet so that  
 627 the stagnation pressure can enter the cavity and cause the  
 628 metal to float and burst. C-bullets expand more rapidly than  
 629 L-bullets and deformation occurs “instantly” when fluid  
 630 pressure enters the anterior cavity. Thereafter penetration  
 631 occurs shoulder stabilized without additional deformation. It  
 632 might happen, though, that petals are lost if heavy bones are  
 633 penetrated (Kneubuehl et al. 2011). L-bullets on the other  
 634 hand will be more liable to change their anterior profile after  
 635 initial expansion because they will be influenced as long as  
 636 the stagnation pressure exceeds the yield limit for lead  
 637 (MacPherson 1994). This probably contributes to a greater  
 638 variability of the anterior surface for the retrieved L-bullets  
 639 compared to C-bullets. Thus, C-bullets exhibited a more  
 640 symmetrical deformation history. One advantage of a sym-  
 641 metric anterior leading surface should be less deviation from  
 642 a strait propagation line in tissue compared to L-bullets with  
 643 a higher degree of asymmetrical expansion (Kneubuehl et al.  
 644 2011). Heterogenous tissues might, however, cause any  
 645 bullet to deviate substantially from a straight line (Kneu-  
 646 buehl et al. 2011).

647 Some concern has been raised regarding the potential of  
 648 C-bullets to expand at longer ranges (Caudell et al. 2012;  
 649 Caudell 2013). This is because loss of flight speed due to  
 650 drag will reduce fluid pressure in the frontal cavity of the  
 651 bullet at impact so that expansion will be reduced or fail to  
 652 happen et al. Within shooting distances applied in the  
 653 present study (150 m), we did not detect any sign of such  
 654 malfunction. Therefore, we did not include shooting dis-  
 655 tance as a factor influencing killing performance in our  
 656 approach. Another interesting observation was that relative  
 657 expansion of bullets decreases with increasing caliber.  
 658 With increasing caliber, ballistic velocity decreases,  
 659 whereas the amount of metal increases (lead or copper).  
 660 This means that there is more metal mass to move during  
 661 expansion when the stagnation pressure forces the metal to  
 662 float. As a result, relatively less metal is probably shuffled  
 663 during expansion resulting in reduced relative expansion.

664 Since the primary factor causing permanent wound  
 665 cavity in soft tissues, such as lungs, mainly is crushing  
 666 rather than radial stretching (Stefanopoulos et al. 2014),  
 667 there should be a correlation between the radial dimension

of the permanent wound cavity and expansion indices. The  
 expansion advantage (larger indices) we registered for  
 C-bullets apparently enables them to compensate for the  
 efficiency of fragmenting so typical for L- bullets. Tri-  
 nogga et al. (2013) also found that permanent wound  
 cavities caused by deforming copper bullets tended to be  
 the largest of all bullet types.

Even though our study indicates that there is no con-  
 sistent and significant difference between the efficacy of L-  
 and C-bullets for hunting, we will suggest one possible way  
 for further improvement of the present incapacitation  
 power of C-bullets. Based on our experiences with the  
 present study, one way of improving the incapacitation  
 power of C-bullets is to increase the expansion index by  
 increasing the ability to expand. Energy transfer strongly  
 depends on the size of the frontal area of the expanded  
 bullet (Wolberg 1991). Therefore, penetration depth  
 decreases as bullet expansion increases. So, the question is,  
 will C-bullets manage to retain their penetration ability in  
 combination with increased expansion? Much of the  
 rationale behind the development of C-bullets was to  
 improve bullet mass retention during expansion to maxi-  
 mize the ability of penetration and wounding (Thomas  
 et al. 2016). However, we did not detect any significant  
 difference between the two bullet types regarding pene-  
 tration ability, so it might be that this ability will restrain  
 further development of expansion indices for C-bullets.  
 The problem might be omitted by making C-bullets heav-  
 ier. Such a solution may, however, cause problems because  
 it implies increased bullet length making them more liable  
 to lose stability, because the distance between center of  
 gravity and center of pressure (air drag) increases (Carlucci  
 and Jacobson 2014). This applies especially to the smallest  
 calibers (i.e., 6.5 mm and smaller) where barrel twist is  
 insufficient to stabilize longer bullets.

## CONCLUSIONS

We found no appreciable difference in killing efficiency  
 between copper and lead-based bullets in our study, which  
 was based on data collected by hunters under normal  
 hunting conditions in Fennoscandia. We evaluated the  
 efficiency of copper versus lead-based ammunition in  
 relation to a quantifiable animal welfare standard. We did  
 not detect any significant difference between reported  
 animal flight distances between copper and lead-based  
 ammunition relative to our standardized predicted animal  
 flight distances based on body mass. Copper ammunition  
 exhibited a larger, more reliable and stable expansion  
 compared to lead-based ammunition. This characteristic  
 seems to offset the advantage lead-based ammunition has  
 in terms of killing efficiency due to fragmentation effects.

718 Given the considerable documentation of harmful health  
719 and environmental impacts from lead-based ammunition,  
720 hunters should strongly consider using copper-based  
721 ammunition based on the results of our study.

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732 provided data on hunting practices.  
733

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