Comparative changes in density and demography of large herbivores in the Masai Mara Reserve and its surrounding human-dominated pastoral ranches in Kenya

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Abstract  Wildlife habitats in pastoral lands adjoining protected areas in east African savannas are getting progressively degraded, fragmented and compressed by expanding human populations and intensification of land use. To understand the consequences of these influences on wildlife populations, we contrasted the density and demography of 13 wild and three domestic large herbivores between the Masai Mara National Reserve and the adjoining pastoral ranches using aerial surveys conducted in the wet and dry seasons during 1977–2010. Species of different body sizes and feeding styles had different densities between landscapes and seasons. Small-sized herbivores, requiring short, nutritious grasses, and browsers were more abundant in the ranches than the reserve in both seasons.

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Medium-sized herbivores moved seasonally between landscapes. Larger-bodied herbivores, requiring bulk forage but less susceptible to predation, were more abundant in the reserve than the ranches. The proportions of newborn warthog (*Phacochoerus africanus*) and juvenile topi (*Damaliscus korrigum*) were higher in the ranches, with shorter grasses and lower predation risk than in the reserve. These results suggest that pastoral lands adjoining protected areas in African savannas are important as seasonal dispersal and breeding grounds for wild herbivores. However, human population growth and dramatic land use changes are progressively degrading wildlife habitats in pastoral areas, thus restricting the seasonal wildlife dispersal movements between the protected areas and adjoining pastoral lands. Conservation efforts should focus on (1) creating and maintaining functional heterogeneity in protected areas that mimic moderate pastoral grazing conditions to attract small and medium-bodied grazers and (2) securing dispersal areas, including corridors, to ensure continued seasonal large herbivore movements between protected and pastoral systems.

**Keywords** Large herbivores · Pastoralism · Protection · Distribution · Seasonal movements

**Introduction**

Pastoralism is the economic mainstay of most inhabitants of grasslands of East Africa, who also often derive limited income from wildlife-based tourism. However, rapid human population growth, expansion of settlements (Lamprey and Reid 2004), cultivation (Serneels et al. 2001; Thompson and Homewood 2002) and transition from semi-nomadic pastoralism to a sedentary lifestyle (Western et al. 2009), are progressively altering the vegetation composition and structure of these savanna grasslands. Concurrent with these processes, a transition from communal land tenure to private land ownership in the pastoral ranches, habitat fragmentation through land privatization and subsequent subdivision (Galvin et al. 2008; Homewood et al. 2009), rising temperatures and recurrent severe droughts (Ogutu et al. 2007) threaten the future survival of large mammalian populations in some savanna ecosystems, such as the Mara-Serengeti of Kenya and Tanzania (Ottichilo et al. 2001; Ogutu et al. 2009).

Settlements are expanding faster nearer than farther away from protected areas in Latin America and Africa due to enhanced economic activities and opportunities inside and around protected-area boundaries (Wittemyer et al. 2008). A spectacular example of this expansion is found on pastoral ranches surrounding the Masai Mara National Reserve (MMNR) in Kenya (Norton-Griffiths et al. 2008). The progressive intensification of land use, sedentarization and diversification of livelihoods are associated with rapidly declining wildlife numbers in the last three decades in pastoral systems of east Africa, including the Mara (Brotten and Said 1995; Ottichilo et al. 2000; Ogutu et al. 2009), Laikipia (Georgiadis et al. 2007) and Athi-Kaputiei (Reid et al. 2008) regions of Kenya and the Tanzanian Tarangire-Simanjiro Plains (Msoffe et al. 2011). The declines are related to increasing numbers of settlements, people, poaching and major land use changes on the pastoral ranches (Serneels and Lambin 2001; Georgiadis et al. 2007; Reid et al. 2008; Ogutu et al. 2009). The patterns of declining wildlife in protected areas of East Africa (Stoner et al. 2007; Western et al. 2009) are consistent with early forecasts of major reductions, and even extinctions of many wildlife populations expected in East African reserves as a
consequence of increasing insularization (Newmark 1996) and displacement of wildlife by increasing livestock incursions into protected areas (Butt et al. 2009).

These changes progressively impede traditional seasonal wildlife movements between protected areas and their adjoining pastoral systems. Several studies have demonstrated seasonal movements by ungulates between protected areas and adjoining pastoral ranches in Amboseli (Western 1975; Mworia et al. 2008), Mara (Stelfox et al. 1986) and Athi-Kaputiei Plains (Reid et al. 2008), thus supporting the prediction that the processes associated with land use change will continue to erode grazing areas so that livestock will compete increasingly with wildlife for resources, resulting in wildlife and livestock population declines (Homewood et al. 2009).

By moving seasonally between protected and pastoral areas, ungulates maximize their resource requirements while minimizing predation risk (Hopcraft et al. 2010). However, these seasonal dispersal movements might be constrained by body size (Hopcraft et al. 2011) through its influence on food quantity and quality requirements as well as vulnerability to predation.

More specifically, large herbivores can tolerate more fibrous and lower-quality diets than can small herbivores because of their larger gastrointestinal tracts and lower specific metabolic requirements (Demment and Van Soest 1985; Owen-Smith 1988). Furthermore, a smaller fraction of large herbivores die from predation than do small herbivores because large herbivores are more difficult for predators to capture (Sinclair et al. 2003). Thus, body size can be expected to control responses of herbivore abundance to seasonal disparities in forage quantity and quality and predation risk between protected and pastoral landscapes.

The MMNR in Kenya supports a high abundance and diversity of resident wildlife and offers a dry season habitat for migratory ungulates from the Serengeti National Park in Tanzania to the south and the neighbouring Loita Plains to the northeast (Stelfox et al. 1986; Ottichilo et al. 2001; Thirgood et al. 2004). Extensive grasslands in the pastoral areas adjacent to the MMNR also provide wet season dispersal ranges for resident wildlife (Stelfox et al. 1986). Yet, despite the significance of pastoral areas to wildlife, few studies have evaluated the relative impact of pastoralism versus protection on wildlife population density and demography in African savannas (Caro 1999a; Rannestad et al. 2006; Wallgren et al. 2009). Even fewer studies have investigated the impacts of pastoralism and protection on long-term comparative changes in density (Caro 1999b; Reid et al. 2008).

Here, we analyze the influence of protection in the MMNR and pastoralism in the adjoining Koyiaki pastoral ranch (see below) on comparative changes in the density of 13 wild herbivores. We consider the following wild herbivores, in order of increasing body size, Thomson’s gazelle (*Gazella thomsonii*), impala (*Aepyceros melampus*), warthog, Grant’s gazelle (*Gazella granti*), topi, wildebeest (*Connochaetes taurinus*), Coke’s hartebeest (*Alcelaphus buselaphus cokii*), defassa waterbuck (*Kobus ellipsiprymnus*), zebra (*Equus burchelli*), eland (*Taurotragus oryx*), buffalos (*Syncerus caffer*), giraffe (*Giraffa camelopardalis*), and elephant (*Loxodonta africana*) (Table 1). We also consider the densities of three domestic herbivore species, namely sheep (*Ovis aries*), goats (*Capra hircus*) and cattle (*Bos indicus*). We used data collected from systematic reconnaissance aerial surveys conducted during wet and dry seasons by the Kenya Department of Resource Surveys and Remote Sensing (DRSRS) from 1977 to 2010. We supplemented these comparisons with parallel comparisons based on ground mapping censuses conducted in the MMNR and Koyiaki in November 1999 and 2002 (Reid et al. 2003). We also compared age and sex composition counts of a subset of six of the 13 wild herbivores, namely, impala, warthog, topi, hartebeest, zebra and giraffe, conducted in 2003 and 2004 to establish the influence of protection and pastoralism on the demography of these herbivore...
species. The six species were selected because reliable methods for ageing and sexing them had already been developed and tested as part of a 15-year monitoring program spanning 1989–2003 (Ogutu et al. 2008).

Our hypotheses were based on differences in grass heights and predator densities between the MMNR and the pastoral ranches quantified by Ogutu et al. (2005) and Reid et al. (2003). Grass height influences both forage quality and predation risk. In the wet season less heavily grazed grasses, such as occur in most parts of the Mara reserve, become tall and therefore allocate more energy to developing structural fibers with higher carbon to nitrogen ratios, thereby diluting the concentration of nitrogen and phosphorous available to herbivores (Anderson et al. 2007). From an herbivore’s perspective, the digestibility of grasses is therefore inversely related to rainfall amount (Hopcraft et al. 2011). Mature grasses of tall stature are thus particularly unfavourable for small and medium herbivores due to their low digestibility and nutritional quality (Fritz and Duncan 1994; Olff et al. 2002). In contrast, short grasses maintained by heavy livestock grazing, such as those in the pastoral areas of the Mara in the wet season (Ogutu et al. 2005), have higher digestibility and nutritional quality. Heavy livestock grazing on the ranches, furthermore, tends to promote production of more net grass biomass, which in turn attracts more herbivores than in the reserve with no livestock. Consequently, sustained livestock grazing in the ranches, by keeping grass stem biomass low, renders grasses more digestible and enhances their nutritional quality (McNaughton 1976). This enables herbivores to realize greater protein consumption on the ranches than they do in the reserve in the wet season. As well, nutrient-rich pastoral settlement (boma) sites in the ranches represent key sources of nutritionally sufficient forage, especially for lactating females in the wet season (Muchiru et al. 2008; Augustine et al. 2010).

In addition, during the wet season, it is likely that lions are more abundant in the reserve (Reid et al. 2003), with taller grass cover, than in the ranches (Ogutu et al. 2005). Predator

| Common name        | Scientific name             | Mass (kg) | Dietary guild | Residence guild |
|---------------------|-----------------------------|-----------|---------------|-----------------|
| Thomson’s gazelle   | *Gazella thomsoni*          | 15        | Grazer        | Migratory       |
| Sheep + goats       | *Ovis aries* + *Capra hircus* | 16        | Mixed feeder^b| Resident        |
| Impala              | *Aepyceros melampus*        | 40        | Mixed feeder  | Resident        |
| Warthog             | *Phacocoerus africanus*     | 45        | Grazer        | Resident        |
| Grant’s gazelle     | *Gazella granti*            | 50        | Mixed feeder  | Resident        |
| Topi                | *Damaliscus korrigum*       | 100       | Grazer        | Resident        |
| Wildebeest          | *Connochaetes taurinus*     | 120       | Grazer        | Migratory       |
| Hartebeest          | *Acelaphus buselaphus cokes*| 125       | Grazer        | Resident        |
| Defassa waterbuck   | *Kobus ellipsiprymnus*      | 160       | Grazer        | Resident        |
| Cattle              | *Bos indicus*               | 180       | Grazer        | Resident        |
| Zebra               | *Equus burchelli*           | 200       | Grazer        | Migratory       |
| Eland               | *Taurotragus oryx*          | 350       | Mixed feeder  | Migratory       |
| Buffalo             | *Syncerus caffer*           | 700       | Grazer        | Resident        |
| Giraffe             | *Giraffa camelopardalis*    | 1,250     | Browser       | Resident        |
| Elephant            | *Loxodonta africana*        | 5,500     | Mixed feeder  | Dispersal^a     |

^a Wanders widely seasonally but do not engage in regular seasonal migrations

^b Sheep are grazers, and goats are browsers
densities are also higher in the reserve than in the ranches in the dry season (Reid et al. 2003), reflecting not only their preference for high grass cover, but also avoidance of human and livestock activities on the ranches (Ogutu et al. 2005). Since predation risk increases with grass height in the Serengeti (Hopcraft et al. 2005) and Mara Region (Kanga et al. 2011) and since grass cover is shorter and predator density is lower on the ranches than in the reserve, small and medium herbivores likely experience lower predation risk on the ranches than in the reserve (Sinclair et al. 2003).

In the dry season, when surface water and forage availability are reduced, heavy livestock grazing in the pastoral ranches forces wildlife to disperse to the reserve, where the migratory wildebeest and zebra and fires have removed the taller grasses and improved visibility. Thus, heavy livestock grazing in the pastoral ranches facilitates small and medium-sized herbivores in the wet season, but competition with livestock in the dry season for food and water, pushes them into the reserve where they are facilitated by migratory herds, which also absorb most of the predation pressure (Ogutu et al. 2008).

Accordingly, we formulated the following four initial expectations based on herbivore body size. (1) The densities of the small-sized herbivores (15–50 kg), would be higher in the Koyiaki pastoral ranch in both seasons due to the higher prevalence of short grass that is safer year round. (2) The densities of medium-sized grazers (100–200 kg) would be higher in the Koyiaki ranch in the wet season when grass is short (and safe) and green (and nutritious), but would be higher in the reserve with higher quantities of grass in the dry season when grass dries out and becomes too short on the Koyiaki ranch. (3) The density of large wild herbivores (>350 kg) would be higher year-round in the reserve than in Koyiaki ranch if they perceive lower predation risk (Sinclair et al. 2003) and satisfy their energy demands by ingesting large quantities of low-quality forage (Demment and Van Soest 1985). Finally, (4) the lower number of predators and presumably lower predation risk on Koyiaki ranch, due to the shorter grasses of higher nutritional quality, and better predator visibility, would lead to a higher proportion of the pregnant females bearing and raising their young on the ranches than in the reserve.

Since the changes in wildlife distribution between the reserve and the ranches constitute essentially an unreplicated natural experiment, we used the protected Mara reserve as an ecological baseline area or benchmark that is relatively free of human impact to understand the consequences of impacts of livestock and human use of the human-dominated pastoral lands on seasonal and long-term patterns of wildlife distributions in the Mara Region (Sinclair 1998; Sinclair et al. 2002). We conduct replicate comparisons of herbivore densities between the reserve and the ranches based on 50 independent aerial surveys spanning 41 years conducted using the same technique to increase our confidence in, and ability to, separate the impacts of livestock and human use of the pastoral ranches on wildlife distributions despite the lack of true replication, which is difficult to achieve experimentally at landscape scales.

Study area

The Mara Reserve is located in southwestern Kenya and borders the Serengeti National Park in Tanzania to the south. It covers some 1,530 km² and is bounded by the Siria escarpment on the west, Koyiaki (931 km²) and Olkinyei (804 km²) pastoral ranches on the north and Siana pastoral ranch (1,315 km²) on the east (Ogutu et al. 2005) (Fig. 1). The reserve and the surrounding pastoral areas support annual migrations of enormous herds of wildebeest and zebra and small herds of eland from the Tanzanian Serengeti and much smaller herds of wildebeest, zebra and Thomson’s gazelles from the Kenyan Loita Plains,

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to the northeast of the reserve (Maddock 1979; Stelfox et al. 1986). Traditional pastoralism, cultivation, and wildlife tourism constitute the major forms of land use in the pastoral ranches (Homewood et al. 2001). The major livestock species kept in the ranches include cattle, sheep, goats and donkeys (Lamprey and Reid 2004). The reserve is a nationally protected area in which wildlife conservation and tourism are the only permitted land uses but illegal livestock grazing is common, especially in dry years (Reid et al. 2003; Butt et al. 2009). There is no physical barrier to wildlife movements between the reserve and the surrounding pastoral areas. Hereafter, we refer to the reserve and all its surrounding pastoral ranches as the “Mara Region”.

The vegetation consists predominantly of open grass plains dominated by *Hyparrhenia filipendula* and *Themeda triandra*, interspersed with *Acacia gerrardii* and *Terminalia* trees, shrublands and riverine forests (Epp and Agatsiva 1980). Numerous seasonal streams drain the area, but only the Mara River and sections of the Sand and Talek Rivers typically contain water year-round. The Mara River originates in the Mau escarpment to the north of the Mara region. Annual rainfall during 1989–2003 averaged 1,010 mm and increased from 877 mm at Ololaimutia Gate in the southeast to 1,341 mm at Kichwa Tembo in the northwest of the MMNR (Ogutu et al. 2011). Rainfall is bimodal in the Mara Region, with the wet season spanning late November of the previous year to June of the current year and the dry season covering July-early November of the current year. The short rains fall during late November–December and the long rains during March-June. Rainfall increases spatially from 500 mm per year in the Serengeti Plains in the southeast to over 1,200 mm in the northwest of the Mara Region (Pennycuick and Norton-Griffiths 1976).
Methods

The Kenya Department of Resource Surveys and Remote Sensing (DRSRS) conducted 50 aerial surveys in the Mara Region from 1977 to 2010, covering the entire Mara Region (6,400 km²), including the reserve (1,530 km²), and the surrounding pastoral ranches (4,870 km²). Surveys were undertaken either in the wet (Jan–June or Nov–Dec) or dry (Jul–Oct) season month(s) of each year except 1981, 1988, 1995, 1998, 2001, 2003, 2004 and 2006 when surveys were not conducted due to financial constraints (Stelfox et al. 1986; Broten and Said 1995; Ottichilo et al. 2000, 2001; Ogutu et al. 2011). The surveys followed systematic strip transects located 5 km apart and segmented into sampling grid cells of $5 \times 5$ km² (Norton-Griffiths 1978). The transects were oriented in an east–west or north–south direction and were flown at a fixed height of about 90 m above the ground during 1977–1985 and about 120 m thereafter (Ottichilo et al. 2000). The number of animals observed within a calibrated survey strip defined by two parallel rods on the wing struts of the aircraft and running through the centre of the $5 \times 5$ km² grid cell was recorded. The survey strip spanned an average width of 263 m on the ground, corresponding to an average sampling intensity or fraction of 4.8% of the $5 \times 5$ km² grid cell area (Ogutu et al. 2011). The expected number of animals per 25 km² grid cell area was thus estimated as the actual number counted in each 25 km² grid cell times 100 divided by the sampling fraction. The mean count for each species per survey in the reserve was expressed as the average of the estimated population size over all the 25 km² grid cells in the reserve ($n = 61$ cells, covering a total area of 1,525 km²). The same applies to the Koyiaki pastoral ranch ($n = 37$, for a total area of 925 km²). Ottichilo (1999) and Ottichilo and Khaemba (2001) have demonstrated the reliability of the estimates of wildlife and livestock population sizes from the DRSRS count method. From the 50 surveys, we selected counts of 13 wild herbivore species, comprising four small-sized herbivores: Thomson’s gazelle, Grant’s gazelle, impala and warthog, five medium-sized herbivores: topi, hartebeest, wildebeest and zebra, four large herbivores: eland, buffalo, giraffe and elephant; and three species of livestock, namely sheep and goats (which are lumped together during surveys as ‘sheep and goats’ because they occur in mixed herds that are hard to distinguish reliably from the air) and cattle to represent a range of functional groups based on body size, feeding and foraging styles (Table 1). Of the 50 surveys 33 were conducted in the wet season and 17 in the dry season. Averaging population density estimates for each species in each grid cell over all surveys conducted in each season in 1 year produced 20 surveys for the wet season (late November–June) and 12 for the dry season (July-early November), which we used for analysis.

Ground mapping census of wildlife and livestock

Two ground mapping censuses of wildlife and livestock in the MMNR and the adjacent pastoral ranches were conducted in early November 1999 and 2002 when dry conditions prevailed and the grass was still short, due to heavy grazing by migratory wildlife (Reid et al. 2003). The first census covered an area of 1,544.2 km², including sections of Koyiaki and Lemek pastoral ranches, and the MMNR. This census was carried out by 12 teams totaling 40 people using 12 vehicles in both the reserve and the ranches. The second census covered 2,212 km² and included Koyiaki, Lemek, Siana and a small part of southwestern Olkinyei ranches. This census was carried out by 22 teams totaling 84 people. The census area was partitioned into contiguous $0.33 \times 0.33$ km² sub-blocks to obtain fine resolution counts. The teams counted 7,606 sub-blocks in the reserve and 6,295 sub-blocks in the
ranches in 1,999 and 11,117 sub-blocks in the reserve and 8,794 sub-blocks in the ranches in 2002 (Reid et al. 2003; Ogutu et al. 2010). The sampling teams navigated vehicles down the centers of each 1 × 1 km² block and allocated all animals observed into one of the nine nearest 0.33 × 0.33 km² sub-blocks using a global positioning system (GPS). The counts per 0.33 × 0.33 km² sub-blocks were converted to densities per km² by multiplying them by nine. The mean density and corresponding standard errors were calculated as the average density over all sub-blocks in the reserve and ranches. The mean count for each species in the reserve was expressed as the average of the estimated population size over all the per 0.33 × 0.33 km² sub-blocks in the reserve. The same applies to Koyiaki pastoral ranch. We used these censuses to validate distribution patterns derived from the aerial surveys during the dry season, including one aerial survey that was conducted at the same time as the ground mapping census in 2002.

Age and sex composition counts of wildlife

Ogutu et al. (2006), in collaboration with the World Wide Fund for Nature (WWF), carried out two further vehicle ground sample counts of impala, warthog, topi, hartebeest, zebra, and giraffe including their age and sex. These counts were conducted in the MMNR, Koyiaki and a small section of Siana ranch in November 2003 and April 2004. The November 2003 survey was also conducted during the dry season. In contrast, the April 2004 survey was conducted in the late-wet season. They used a strip-transect sampling technique assuming complete census of all animals within a fixed strip width of 100 m either side of the transect centerline (Ogutu et al. 2006). The transects were distributed over the MMNR and pastoral ranches in proportion to their areas, with 22 transects established in the reserve and 13 in Koyiaki. Each transect was 10 km long. After every 1 km along each transect, the vehicle was stopped and the numbers, age class relative to adult size, sex and GPS locations of wildlife were recorded within 200 m on either side of the transect centerline. These species were classified, whenever possible, into three age classes: newborns (<1 month), juveniles (1–18 months), adults (>18 months). A combination of horn shape and length and body size were used to assign the herbivores to sex and age categories, however, ages were not assigned to adults (Sinclair 1995; Ogutu et al. 2008). Only the number of individuals sighted per age class in each transect, summed over all transects in the reserve and the ranches, from this dataset were used in analyses.

Comparing wildlife and livestock densities between landscapes

To account for clustering, non-normality and non-homogenous variances of animal counts, and varying frequency of counts we used negative binomial regression model for over-dispersed count data to compare the mean density for each herbivore species in each 5 × 5 km² grid cell between the MMNR and Koyiaki pastoral ranch using the aod package in R (Lesnoff and Lancelot 2010; R Development Core Team 2010). More specifically, we used the log link function and specified the variance function for the negative binomial model as \( \varphi u(1 + (u/k)) \), where \( u \) is the mean, \( \varphi \) is the overdispersion parameter and \( k \) is the ‘aggregation parameter’. Differences in the expected herbivore counts between landscapes were tested for significance using the Wald Chi-squared test (Draper and Smith 1998). A similar analysis was performed to compare the mean densities from the ground mapping censuses per 1 × 1 km² grid cells between the MMNR and Koyiaki pastoral ranch (Reid et al. 2003).
Comparing age ratios and female proportions between landscapes

Due to low sample sizes for certain cross-classification cells, we used a Chi-square test for independence in a $2 \times 2$ frequency table, corrected for continuity using the Yates’ correction for small sample sizes, to compare the ratio of newborns to adult females to that of juveniles to adult females for impala, topi and giraffe and the ratios of newborns to adults of both sexes to that of juveniles to adults of both sexes for warthog and zebra between the two areas based on counts pooled over the 2003 and 2004 surveys. We similarly compared the female proportion ($F/(F + M)$, where $F =$ female counts and $M =$ male counts) for impala, topi and giraffe computed by pooling all individuals of the same sex over all age classes and the 2003 and 2004 surveys, separately for each area.

Results

Comparative changes in herbivore density

The details of differences in wildlife densities between the reserve and the ranches were complex and varied with species and season, but some consistent overall patterns were nevertheless evident.

Small sized herbivores

Most small herbivores were consistently more abundant in the ranches than in the reserve in both seasons (Fig. 2a, e). Interestingly, warthog did not conform to this pattern and showed a preference for the reserve in the dry season but for the ranches in the wet season (Fig. 2d). Sheep and goats were more abundant in the ranches than in the reserve, and their numbers increased noticeably during 2000–2010 relative to earlier years (Fig. 2b; Tables S1, S2).

Medium sized herbivores

Most medium-sized herbivores moved seasonally between the reserve and the ranches (Fig. 3a, f). However, hartebeest and waterbuck had slightly higher densities in the reserve during both seasons, but more especially in the wet season (Fig. 3c, d; Tables S1, S2). Topi, wildebeest and zebra had slightly higher densities in the reserve in the dry season when the migrants are present but somewhat higher densities in the ranches in the wet season (Fig. 3a, b, f; Tables S1, S2). More specifically, the resident wildebeest had lower densities in the ranches than in the reserve in the dry season but higher densities in the ranches than in the reserve in the wet season (Fig. 3b). Cattle were more abundant in the ranches than in the reserve in the dry season but more occurred in the reserve in the dry than in the wet season, and more recently (2000–2010) than in earlier years 1970–1999 (Fig. 3e; Tables S1, S2).

Large sized herbivores

Buffalo and elephant were consistently more abundant in the reserve than in the ranches in both seasons (Fig. 4b, d; Tables S1, S2). Eland had higher densities in the ranches than in the reserve in the wet season but lower densities in the ranches than in the reserve in the dry season (Fig. 4a). Giraffe did not show significant differences between the reserve and the ranches during the dry season, but were somewhat more abundant in the reserve.
However, they were consistently more abundant in the ranches than the reserve in the wet season (Fig. 4c; Tables S1, S2).

The ground counts conducted in 1999 and 2002 confirmed that both gazelles, impala and giraffe were indeed more abundant in the ranches and that topi, hartebeest, wildebeest, zebra, eland, buffalo and elephant were more abundant in the reserve than in the ranches in
the dry season, as revealed by the aerial survey data. High variance in herd sizes rendered the apparently large differences in wildebeest densities between landscapes statistically insignificant. The ground counts also confirmed the greater abundance of livestock in the ranches than in the reserve shown by the aerial survey data (Table 2).

Comparisons of age ratios and female proportions between the reserve and the ranches

The population age composition of species differed between areas for warthog, topi and zebra. There were greater proportions of newborn warthog and juvenile topi in the ranches than in the reserve, but greater proportions of newborn topi and zebra in the reserve than in the ranches (Table 3). For hartebeest and waterbuck, numbers were too small for similar statistical tests. Only impala, topi, hartebeest and giraffe had sufficient sample sizes to statistically test differences in female proportions between the two areas. Among these species, female proportion was similar between landscapes for hartebeest and giraffe but was higher in the reserve than in the ranches among impala and topi (Table 4).

Discussion

Comparative changes in herbivore density

Although the differences in herbivore densities between the pastoral ranches and the reserve in any 1 year may also be influenced by inherent differences between the
landscapes which are unrelated to livestock or human use, such as geomorphology, which can cause underlying differences in wildlife use, our results suggest that livestock and human use of the pastoral ranches are the two most important causes of the differences between the patterns we observed in the two landscapes.

Fig. 3 Comparative changes in densities (number/km²) of medium pure grazers, a topi, b wildebeest, c hartebeest, d waterbuck, e cattle and f zebra between the Mara Reserve (light bars) and the adjoining Koyiaki pastoral ranch (dark bars) during the dry and wet seasons based on the DRSRS aerial surveys from 1977 to 2010. Vertical lines show the 95% pointwise confidence limits whereas stars indicate that the mean densities differed significantly between the reserve and Koyiaki.
There were strong seasonal differences in wild herbivore densities between the reserve and the ranches during 1977–2010. Individual species responded differentially to pastoralism and protection. Three distinct patterns were apparent, all of which could be explained in terms of distinctions in body size and feeding guild and their consequences for nutritional quality and quantity of forage, predation risk and competition with livestock.

Small sized herbivores

Small species that are constrained by food quality and predation tend to prefer short grass areas (Fryxell 1991; Illius and Gordon 1992) and were thus more abundant in the ranches than the reserve regardless of season or feeding guild as revealed by the significant

Fig. 3 continued

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Repeated livestock grazing in the same areas of the ranches probably increased the crude protein production of grasses (Anderson et al. 2010; Augustine et al. 2010), enabling the small grazers to derive sufficient energy by selecting high-quality forage from the low-biomass areas (Fryxell et al. 2005). Reduced predation risk as a result of lower vegetation cover on the ranches (Ogutu et al. 2005) is yet another advantage of concentrating in the short grass plains, since tall grasses conceal ambush predators and significantly increase their efficiency at catching prey animals (Hopcraft et al. 2005). The distribution patterns we observed for small herbivores are therefore concordant with the initial expectation that small herbivores (except warthog) should concentrate in areas of relatively fewer predators (safer) and shorter grasses maintained by heavy livestock grazing in the ranches. This outcome also concurs with findings of studies encompassing a variety of spatial scales and species (Olff et al. 2002; Cromsigt and Olff 2006) besides reinforcing the notion that both predation and resource limitation act simultaneously in limiting herbivore populations (Sinclair et al. 2003).

Medium sized herbivores

The second pattern was expressed by species that moved between the ranches and the reserve seasonally, suggesting that they preferred either the reserve or the ranches depending on season. Specifically, the medium-sized topi, wildebeest and zebra moved seasonally between the reserve and the ranches, thus supporting our second prediction. As a result, medium herbivores had higher densities in the ranches in the wet season but higher densities in the reserve in the dry season. This pattern suggests that medium herbivores tend to utilize the ranches when water and short, nutritious grasses, created and maintained by heavy livestock grazing (Rannestad et al. 2006), are widely available, enabling them to enhance their total protein consumption (McNaughton 1976). In addition, the short grasses also enhance visibility of predators, thus potentially lowering predation risk in the ranches than the reserve (Hopcraft et al. 2005; Ogutu et al. 2005). In contrast, since heavy and sustained livestock grazing depletes both forage and surface water faster in the ranches than in the reserve (Reid et al. 2003), the medium-sized grazers are likely forced to disperse from the ranches to the reserve in the dry season to access more forage and water. Consequently, the medium-sized species were more abundant in the reserve during the dry season, implicating elevated competition with livestock on the ranches for food and water. These patterns accord with the finding of Odadi et al. (2011), who recently reported greater competitive effects of livestock on wildlife in the dry season when food is scarcest.

Interestingly, hartebeest and waterbuck, both medium-sized grazers that select long grasses (Murray and Brown 1993), did not conform to this pattern; instead, they showed a slight preference for the reserve where long grasses are more abundant year-round (Reid et al. 2003; Ogutu et al. 2005). Because zebra can process large quantities of low quality diet due to their non-ruminant digestive physiology than can, say, the ruminant wildebeest (Gwynne and Bell 1968; Ben-Shahar and Coe 1992) it could be argued that zebra should be more abundant in the reserve where tall grasses are more abundant in both seasons (Reid et al. 2003; Ogutu et al. 2005). The occurrence of zebra at high densities in the ranches may
### Table 2: Comparisons of mean herbivore densities between the Mara Reserve (808 km²) and Koyiaki pastoral ranch (649 km²) based on ground mapping censuses conducted in November 1999 and 2002

| Species      | November 1999 | November 2002 |       |       |
|--------------|---------------|---------------|-------|-------|
|              | Ranches       | Reserve       | Ranche| Reserve|
| Thomson’s gazelle | 15.97         | 16.70         | 28.13 | 21.30 |
| Sheep + goats | 31.28         | 2.02          | 61.96 | 9.19  |
| Impala       | 9.24          | 4.49          | 12.22 | 6.08  |
| Warthog      | 0.50          | 0.83          | 0.74  | 1.38  |
| Grant’s gazelle | 1.68          | 1.52          | 1.96  | 2.72  |
| Topi         | 2.68          | 4.38          | 3.79  | 4.21  |
| Wildebeest   | 12.75         | 79.21         | 25.58 | 108.35|
| Hartebeest   | 0.14          | 0.38          | 0.16  | 0.42  |
| Waterbuck    | 0.25          | 0.34          | 0.35  | 0.27  |
| Cattle       | 16.84         | 4.08          | 34.30 | 15.98 |
| Zebra        | 7.90          | 11.95         | 15.80 | 21.01 |
| Eland        | 0.20          | 1.00          | 0.15  | 1.37  |
| Buffalo      | 0.50          | 1.27          | 0.08  | 1.31  |
| Giraffe      | 0.59          | 0.24          | 0.65  | 0.25  |
| Elephant     | 0.07          | 0.56          | 0.09  | 0.55  |

Densities that differ significantly ($P < 0.05$) between the two landscapes in each year are highlighted in bold face font.

### Table 3: Tests for differences in age ratios (newborn/adult females, juveniles/adult females; for warthog and zebra adults of both sexes were used in place of adult females and subadults + adults/total) of each species between the Masai Mara Reserve and Koyiaki pastoral ranch based on pooled data for November 2003 and April 2004

| Species      | Age            | Ranch | Reserve | LCL | UCL | $\chi^2$ | $P$  |
|--------------|----------------|-------|---------|-----|-----|----------|------|
|              |                |       |         |     |     |          |      |
| Warthog      | Newborn        | 0.41  | 0.17    | 0.04| 0.42| 7.58     | <0.01|
| Topi         |                | 0.02  | 0.06    | −0.06|−0.01| 10.44    | <0.01|
| Zebra        |                | 0.004 | 0.02    | −0.02|−0.01| 10.38    | <0.01|
| Impala       | Juveniles      | 0.12  | 0.12    | −0.03|0.02 | 0.10     | 0.74 |
| Warthog      |                | 0.13  | 0.30    | −0.32|−0.01| 3.35     | 0.06 |
| Topi         |                | 0.19  | 0.11    | 0.03| 0.11| 18.10    | <0.01|
| Zebra        |                | 0.07  | 0.08    | −0.03|0.003| 2.23     | 0.13 |
| Giraffe      |                | 0.13  | 0.16    | −0.15|0.09 | 0.06     | 0.79 |
| Impala       | Subadults + Adults | 0.85 | 0.85   | −0.03|0.03 | 0.003    | 0.95 |
| Warthog      |                | 0.45  | 0.52    | −0.28|0.13 | 0.24     | 0.62 |
| Topi         |                | 0.78  | 0.82    | −0.08|0.01 | 2.98     | 0.08 |
| Zebra        |                | 0.92  | 0.59    | 0.01| 0.05| 7.28     | <0.01|
| Hartebeest   |                | 0.81  | 0.78    | −0.16|0.22 | 0.003    | 0.95 |
| Giraffe      |                | 0.79  | 0.74    | −0.10|0.20 | 0.24     | 0.62 |

The total number aged in both landscapes and years was 2,410, 201, 2,284, 175, 7,957, and 183 for impala, warthog, topi, hartebeest, zebra and giraffe, respectively. LCL and UCL are the 95% lower and upper binomial confidence limits for each age ratio, respectively. Bold values indicate the significance assessed at alpha = 0.05.
thus suggest attraction to the short, high-quality grasses there and/or lower predation risk, since zebra suffer heavy lion (*Panthera leo*) predation in the Mara-Serengeti ecosystem (Grange et al. 2004). The short grass plains in the ranches also may provide seasonal predator refugia for lekking topi (Bro-Jørgensen and Durant 2003).

Large sized herbivores

The third pattern involved species that prefer long grasses all year, or for part of the year and, thus are most likely to compete strongly with livestock. These species were more abundant in the reserve than in the ranches. Since species such as buffalo and elephant are exposed to less predation risk because of their very large body sizes (Sinclair et al. 2003), they do not have to avoid areas with high risk of predation (Hopcraft et al. 2011) and can therefore, relatively safely, use areas of high food abundance. Furthermore, by often occurring in large herds these herbivores, reduce predation risk even further. Also, their digestive physiology allows them to utilize the low-quality tall grasses predominantly found inside the reserve to maximize their specific metabolic requirements (Illius and Gordon 1992; Wilmshurst et al. 2000). The distribution patterns of the large herbivores thus conform to the expectation that large herbivores should select areas with taller grasses than small herbivores (Sinclair et al. 2003; Hopcraft et al. 2011). The patterns shown by the large-bodied eland did not conform fully to this expectation. Instead, eland moved seasonally between the reserve and the ranches. It is plausible that short, nutritious forbs which eland selects in the wet season (Watson and Owen-Smith 2000; Augustine et al. 2010) occurred at higher densities in the livestock-dominated areas in the ranches in the wet season. By contrast, giraffe are almost exclusively browsers favouring trees and shrubs and feeding almost entirely on forage at least 1 m off the ground (Owen-Smith and Cooper 1987). The ranches support 11–12% woody cover and the reserve 4% as measured by Reid et al. (2003). This higher abundance of trees and shrubs on the ranches may be partially the result of rocky topography in parts of the ranches, but may also be because combined livestock and wildlife grazing removes more grass fuel on the ranches than in the reserve, thus discouraging extensive fires that suppress tree and shrub establishment (Scholes and Archer 1997). As a result, giraffe were more abundant in the ranches with more trees and shrubs in the wet season.

Comparisons of age ratios and female proportions between landscapes

We predicted that the lower number of predators, lower predation risk, and shorter grass (Ogutu et al. 2005), and better predator visibility (Kanga et al. 2011), will lead to a higher
proportion of the pregnant females bearing and raising their young on the ranches than in the reserve. As expected, newborn warthog and juvenile topi were significantly more abundant in the ranches, suggesting a preference for shorter grass areas where predation risk is lower. Contrary to our expectation, however, the proportions of newborn topi and zebra were higher in the reserve, suggesting a push from pastoralists or a pull by something in the reserve, such as tall and dense grass cover for young to hide. The ratio of females to males varied significantly from parity for impala and topi, for which a female biased sex ratio is common (Sinclair et al. 2000). Our results suggest that female impala and topi were more abundant in the reserve, consistent with our speculation that competition with livestock and disturbance by humans and dogs in the ranches forces more females accompanied by their young into the reserve. Female giraffe and hartebeest were evenly distributed between the reserve and ranches, suggesting little influence of land use on the distribution of females relative to males.

Implications for pastoralism, wildlife management and conservation

Dispersal areas for wildlife in pastoral systems and their adjoining protected areas in African savannas represent wet season refuges for many wild herbivores that range seasonally beyond the protected area boundaries (Ogutu et al. 2008; Augustine et al. 2010).

Our study shows that these areas can, and indeed do, support a high diversity of wildlife, especially in the wet season when resources are widely available due to maintenance of grasslands by livestock in short, nutritious growth stage. However, several other studies have shown that increasing human population growth, settlement, cultivation and sedentarization of formerly semi-nomadic pastoralists in these areas are increasingly restricting seasonal wildlife movements (Serneels and Lambin 2001; Coughenour 2008; Ogutu et al. 2011) and potentially negating their otherwise positive effects on wildlife. These movements give both wildlife and livestock the flexibility and mobility necessary to optimally exploit heterogeneity in resources in space and time, including that caused by the directional impacts of a warming and drying climate (Ogutu et al. 2007). Our results reinforce and extend the conclusions of these studies by also revealing that, even though wildlife evidently move seasonally between the reserve and the ranches, their densities have declined strikingly in both the reserve and the ranches, most likely due to ongoing land use changes (Ogutu et al. 2009, 2011). Land use changes in the pastoral lands thus portend a precarious future for wild herbivores that depend on the pastoral areas. Furthermore, the land use changes exacerbate the adverse effects of recurrent climatic extremes on the availability of forage and water, forcing ever more pastoralists to graze their livestock illegally in protected areas (Butt et al. 2009; Ogutu et al. 2009). The land use changes also likely intensify competition between wildlife and livestock and thus adversely affect demographic processes such as reproduction and juvenile recruitment besides the seasonal dispersal movements of wild herbivores between protected areas and their adjoining pastoral lands. If the ongoing losses of key dispersal areas and calving grounds of wildlife in key ecosystems of East Africa, such as the Mara Region, continue unabated, they will accelerate wildlife population declines (Ogutu et al. 2011) and even cause local population extirpations (Newmark 1996).

We therefore suggest that effective management of pastoral lands as well as their adjoining protected areas in East Africa and possibly elsewhere is urgently necessary and should aim to prevent further losses of wildlife. Furthermore, management should aim to secure dispersal areas, including corridors for seasonal wildlife and livestock movements, and effectively couple traditional knowledge of seasonal herdsmen, management and
scientific knowledge (Reid et al. 2009) into an integrated approach incorporating both protected areas and their adjoining pastoral lands.

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