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# Recent vegetation phenology variability and wild reindeer migration in Hardangervidda plateau (Norway)

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More than others, arctic ecosystems are affected by consequences of global climate changes. The herbivorous plays numerous roles both in Scandinavian natural and cultural landscapes (Forbes et al., 2007). Wild reindeer (*Rangifer tarandus* L.) herds in Hardangervidda plateau (Norway) constitute one of the isolated populations along Fennoscandia mountain range. The study aims to understand temporal and spatial variability of intra- and inter-annual home ranges extent and geophysical properties. We then characterize phenological variability with Corine Land Cover ecological habitat assessment and bi-monthly NDVI index (MODIS 13Q1, 250 m). Thirdly, we test relationships between reindeer's estimated densities and geophysical factors. All along the study, a Python toolbox ("GRiD") has been mounted and refined to fit with biogeographical expectancies. The toolbox let user's choice of inputs and facilitate then the gathering of raster datasets with given spatial extent of clipping and resolution. The grid generation and cells extraction gives one tabular output, allowing then to easily compute complex geostatistical analysis with regular spreadsheets. Results are based on reindeer's home ranges, associated extent (MODIS tile) and spatial resolution (250m). Spatial mismatch of 0.6 % has been found between ecological habitat when comparing raw (100m<sup>2</sup>) and new dataset (250m<sup>2</sup>). Inter-annual home ranges analysis describes differences between inter-seasonal migrations (early spring, end of the summer) and calving or capitalizing times. For intra-annual home ranges, significant correlations have been found between reindeer's estimated densities and both altitudes and phenology. GRiD performance and biogeographical results suggests 1) to enhance geometric accuracy 2) better examine links between estimated densities and NDVI.

Keywords: Raster datasets compiling, biogeography, *Rangifer tarandus* L., home ranges, kernel densities estimation, ecological habitat, phenology

## 1. Introduction

In the information age, « big data » constitutes a large part of the future of ecology and biogeography (Hampton et al., 2013). Transdisciplinary expertise for socio-ecological issues are nowadays relying on multi-factorial and multi-scale analysis for environmental surveys. Massive datasets are easily computed and processed, thanks to tremendous

computer hardware advances and trends for horizontal interactions between experts, citizens and institutions (Kambatla et al., 2014). That said, the era of data-intensive science needs some specific tools to collect and centralize the numerous and heterogeneous datasets related to a studied phenomenon, such as wildlife and herding issues, interacting with cultural and natural systems. Multi-temporal scalar information is another issue, long-term strong trends (climate change, biodiversity loss) interacting with inter-annual variability. Within the scope of environmental impact assessment, and particularly in the field of animal ecology, an equal spatial resolution of biotic and abiotic factors is required to describe, explain and forecast how a population is supposed to move in a limited range of space, according to future bioclimate and biocenoses interactions.

Arctic and subarctic ecosystems are particularly affected by the consequences of global climate change (Uboni et al., 2016). *Rangifer tarandus tarandus* L. constitutes a key-component species, and its role in both natural and cultural sphere is increasingly important. With a wide circumpolar distribution, the capital-breeder and long-distance migratory ungulate moves through taiga and tundra biomes. Its particular geographical distribution and its ecological habitats selection are relevant to ecosystem structure and functioning. The ability of *Rangifer* to cope with its food needs and spatial requirements may represent a good opportunity to increase arctic and subarctic socio-ecosystems resilience to rapid effects of climate change (Post and Pedersen, 2008). At a global scale, populations of wild reindeer are classified as vulnerable by the International Union for Conservation of Nature (IUCN), and currently red-listed by the institution. Warmer winter temperatures, rapid changes in the water cycle, increasing landscape fragmentation and human disturbances contribute to deplete populations, biocenoses communities and individuals (Weladji et al., 2006).

On the Hardangervidda plateau (Norway), the subpopulation of *Rangifer tarandus tarandus* L. is one of the last remaining wild reindeer population in Scandinavia. The population stock has been reduced sevenfold between early 1970s and today (Uboni et al., 2016). With Rondane and Snøhetta sub-populations, those three biogeographical isles could represent a genetic resource contributing to Scandinavian *Rangifer* resiliency to climate change and related cultural and natural landscapes. As such, it is important to improve our understanding of the capital-breeder ungulate migrations during the vegetation growth period and the critical times in its life cycle: response to spring snow melt, calving period and “fat accumulation” period in late summer (Klein, 1990; Skarin, 2008). Our hypothesis are the following: *Rangifer tarandus tarandus* L. has a seasonal behavior, choosing different habitats according to the key moment of its biological cycle during the early spring and the growing season (Skarin, 2008); strong estimated densities of *Rangifer tarandus tarandus* L. depend on the bioclimatic conditions during its migration, and on geographical features such as topography and habitats (Klein, 1990).

Management of merged datasets, here by spatializing relationships between biocenoses and biotopes, may improve GIS toolbox as well as our understanding of *Rangifer tarandus* interactions. The study aims to present a GRiD-toolbox, combining some pre-existent tools in current versions of ArcGIS and QGIS software, applied to a multi-scalar reindeer home ranges study.

## 2. Material and methods

### 2.1 Material

Several free-access datasets, fully described in Table 1, are compiled, merged and analyzed in the study. Downloading datasets is the first step (url access in Table 1). The first basis is the reindeer GPS-tracks database used by Cagnacci et al. (2015) to assess and characterize most European’s largest herbivorous seasonal residence times, associated home ranges and identifying migrational structures. 7 wild female reindeers have been equipped and surveyed using GPS-radio collar. The second dataset is ASTER-GDEM, providing altitudes and other topographical informations (slopes, expositions). The third dataset involves average annual temperatures, interpolated from the Global Historical Climatological Network at high resolution (Hijmans et al., 2005). Corine Land Cover map is the fourth dataset. The Corine program assess ecological habitats and land uses at European scale, with remote sensing and photo-interpretation observations. The downloaded datasets are available for 1990, 2000, 2006 and 2012. In our case,

the 2006 edition is used in order to suit temporally with the locations of wild reindeer. The last dataset is represented by MODIS 13Q1 time series. Bi-monthly temporal acquisition has been chosen, from the beginning of 2000 to the end of 2015. Dates of imagery acquisition is expressed in Julian day. The acquired tiles extent to entire Scandinavia, with a spatial resolution of 250 meters. Since the native dataset is in \*.HDR format, Normalized Difference Vegetation Index (NDVI) band has been chosen, and converted into GeoTiff format.

Different softwares are used in the study, depending on processing steps. First of all, we used GIS software, such as ArcMap 10.2 and Qgis 2.18 for both preprocessing and processing. Then, R-studio and Ade-Habitat-HR (Home Ranges, Callenge, 2006) package allow us to calculate reindeer home-ranges, using Kernel Utilization Density technique, on a bi-monthly basis. Then home ranges have been exported in separate GIS files (\*.SHP). These home ranges and associated quantification of reindeer's densities constitute the key-input for the generation of new datasets. Regular spreadsheets, such as Open Calc 4.1.3 or Office Excel 2016 versions are used all along the study. Both R (package Rcmdr) and Xlstat can be used for complex statistical calculation. For this specific study, Xlstat has been privileged.

**Table 1.** Studied datasets and associated metadata

Dataset type	Reindeer GPS-tracks	DEM	Annual temperature	Ecological habitats	Vegetation phenology
Sampled variable	"KUD_ref_value"	"alti", "expo", "slopes"	"tmoy"	Corine Land Cover habitats	MODIS 13Q1-NDVI, 2007-2009 growing season
Format	*.txt	*.tif	*.bil	*.tif	*.hdr
Data source	<a href="http://datadryad.org/resource/doi:10.5061/dryad.rg0v3">http://datadryad.org/resource/doi:10.5061/dryad.rg0v3</a>	<a href="http://www.eea.europa.eu/">http://www.eea.europa.eu/</a>	<a href="http://www.worldclim.org">www.worldclim.org</a>	<a href="http://www.eea.europa.eu/">http://www.eea.europa.eu/</a>	<a href="https://earthexplorer.usgs.gov/">https://earthexplorer.usgs.gov/</a>
Period available	3/20/2007-2/15/2010	-	1960-1990	2006	2000-2015
Geographical extent	Hardangervidda plateau (Norway); up to 100 km <sup>2</sup>	World, tiles	World	Europe	World, tiles
Temporal resolution	3 hours	-	Annual	1990-2000-2006-2012	Bi-monthly
Spatial resolution	~meters	30 meters	900 meters	100 meters	250 meters
Unit of measurement/type of variable	-	Meters, degrees	Celsius	Categories	NDVI index
Geodesic system	WGS84-UTM 32N	ETRS 89	WGS 84	ETRS 89	Polar sinusoidal

## 2.2 Methods: defining spatial units, extracting values and statistical analysis

The general approach requires to define the spatial and temporal unit to compare and to analyze, in this case bi-monthly home ranges during the growing season (April to August). Grid extent will be generated at the regional scale and related cells used for the extraction will depend on bi-monthly home ranges distribution. The spatial and temporal resolution is forced here on the MODIS-13Q1 accuracy, but this can be specified in GRiD-toolbox parameters.

### 2.2.1 Monitoring wild reindeer (*Rangifer tarandus* L.) home ranges in Hardangervidda Plateau (Norway)



### 2.2.3.1 Checking correspondences between raw and sampled datasets

We quantify the possibility to lose spatial information, by upscaling geophysical datasets at MODIS – 13Q1 spatial resolution (~250m). In the study, the comparison will be made between raw Corine Land Cover dataset (e.g. 100m of spatial resolution) and new Corine dataset extracted using GRiD toolbox (250m). The analysis involves the calculation of particular and total error (%) between raw and new dataset. A threshold of 5% of total error has been retained.

### 2.2.3.2 Removing geophysical factors which are spatially autocorrelated

The goal is here about deleting geophysical factors which could be redundant in the analysis between reindeer's home ranges and reindeer's kernel densities within bi-monthly home ranges. At the scale of one of the most extent home range (2<sup>nd</sup> half of April), continuous values (annual temperatures, altitudes, exposition, slopes, NDVI) are crossed using Pearson's correlation. A threshold of 0.8 is retained for the significance of a relationship, allowing then to remove redundant factors.

## 2.2.4 Statistical processing at intra-annual scale of analysis

### 2.2.4.1 Characterizing the seasonal variability of the chosen ecological habitats

For the characterization of the inter-seasonal variability of the choice of ecological habitats, we summed every pixel belonging to each habitat class within the Kernel Utilization Distribution area. Adding up pixels allow us to compare habitat's areas (sum of pixel's absolute values, or percentage calculation) between seasonal home ranges, aggregated during the whole period. The statistical summaries and bar charts for every home range are then summed up by selecting the first and the second most frequent ecological habitat by bi-monthly reindeer's home range. A Khi<sup>2</sup>-test is then computed to verify the independence between the fortnight and the category of land cover (number of pixels).

### 2.2.4.2 Inter-annual variability of bioclimatic conditions: choice of particularly different years for spring migration, calving area and "fat accumulation" area

When extracting MODIS-NDVI time series for annual home ranges, care was taken to sample the three Julian date acquisition relevant to bi-monthly home range. It gives three NDVI time series (columns) by bi-monthly home range. For example, when extracting geophysical raster based on 1<sup>st</sup> half of April, the 97<sup>th</sup> Julian day of 2007, 2008 and 2009 have been added to the 1<sup>st</sup> half of April home range. We then check the independency of these three inter-annual statistical distribution, using Kruskal-Wallis independency tests. Based upon the median of rank generated distributions, two different years in regards of NDVI inter-annual distributions have been selected.

## 2.2.5 Statistical processing at inter-annual scale: specific bi-monthly home ranges

### 2.2.5.1 Testing relationships between biotic and abiotic factors

The selected contextual bi-monthly home ranges correspond to three major moment in reindeer's annual biology (descent to summer pastures = 1<sup>st</sup> half of April, calving time = 2<sup>nd</sup> half of May, fat accumulation time = 2<sup>nd</sup> half of July). For those particular home ranges, we tested the relationships between contextual estimated densities and other

extracted factors (altitudes, exposition, slopes or proxies of primary chlorophyll production such as NDVI). Spearman correlation test is preferred in this case. According to Spearman table of significance, with a confidence threshold of 95% and a number of observations >100, the given threshold for significance is above 0.197.

### 2.2.5.2 Differences in habitat choices

We compare the NDVI values distribution and the reindeer's densities probabilities distributions belonging to the same home range. Our aim is to verify the similarities or dissimilarities between these two kinds of distributions. Kruskal-Wallis independence tests has been conducted using habitat categories as sub-samples. Ecological habitat categories divide both densities estimation and NDVI values as many sub-samples as ecological classes the observations belong to.

## 3. Results and suggestions

### 3.1 GRiD toolbox: sufficient accuracy

**Table 2.** Accumulated error, in percentages, between raw Corine Land Cover dataset (spatial resolution = 100 m<sup>2</sup>) and extracted Corine Land Cover using GRiD toolbox (spatial resolution = 250 m<sup>2</sup>) for the 2<sup>nd</sup> half of April

% of total area for each present class of habitat for raw Corine Land Cover dataset															
Dis-continuous urban fabric (2)	Sport and leisure facilities (11)	Non-irrigated arable land (12)	Complex cultivation patterns (20)	Land principally occupied by agriculture (21)	Broad-leaved forest (23)	Coniferous forest (24)	Mixed forest (25)	Moors and heathland (27)	Transitional woodland shrub (29)	Beaches, dunes, sands (31)	Sparsely vegetated areas (32)	Peat bogs (36)	Water courses (40)	Water bodies (41)	Total
0.04	0.05	0.07	0.30	0.77	16.51	9.49	5.80	38.96	0.55	0.24	15.14	5.45	0.02	6.60	100
% of total area for each present class of habitat, extracted Corine Land Cover dataset, using GRiD															
0.04	0.04	0.08	0.32	0.75	16.40	9.41	5.84	39.16	0.57	0.24	15.14	5.44	0.04	6.54	100
Accumulated error for each present class of habitat and total of overall accumulated error															
0	0.01	0.01	0.01	0.02	0.11	0.08	0.03	0.2	0.03	0	0	0	0.02	0.07	0.6

Table 2 compares the percentages of Corine Land Cover ecological habitats (1<sup>st</sup> line, table 2) according to the spatial resolution, in one of the most extent home range (2<sup>nd</sup> half of April, e.g. 17236 cells; ~4309 km<sup>2</sup>). Similar percentages can be found within coarse dataset (3<sup>rd</sup> line) compared to the percentages for GRiD-extracted dataset. The last line subtracts the difference between GRiD-extracted percentages to coarse Corine Land Cover. Each accounted error has been added up to the total, giving a total error about 0.59 %. Close to 99.4% of representativeness, the GRiD-toolbox extraction shows good results, in particular for Corine Land Cover ecological habitats.

That said, some improvements could reinforce spatial accuracy of GRiD-toolbox. Firstly, the quantification of error between specific spatial resolution of coarse datasets and inputted spatial resolution using GRiD-toolbox has to be more thoroughly assessed. It could be interesting to repeat the same test for other datasets, such as raw ASTER-GDEM

(spatial resolution e.g. 30m) and extracted values (250m). At the scale of the analysis (here the 2<sup>nd</sup> most extent reindeer's home range), it is important to notice that the number of observations (e.g. pixels) can vary according to the considered spatial unit. Since the number of geostatistical observations is very different between home ranges, checking spatial unit representativeness for each dataset would allow us to have a good overview of total error for a given spatial resolution as GRiD toolbox input parameter.

The second improvement is related to geometrical bias, probably generated by the chosen map projection (UTM-32N). Indeed, Mercator projection is more suited with equatorial region, and tends to elongate surfaces when dealing with polar or subpolar regions. Working with an official map projection clipped by zone (EUREF89-NTM in our case) will likely improve overall results for GRiD-toolbox and wild reindeer distribution analysis. Another parameter (choosing cartographic projection) could be added to the toolbox and automatically process the projection homogenization between the numerous datasets we are willing to extract.

### ***3.2 Intra-annual scale: towards a selection of biotic and abiotic parameters***

#### **3.2.1 Auto-correlated and redundant data**

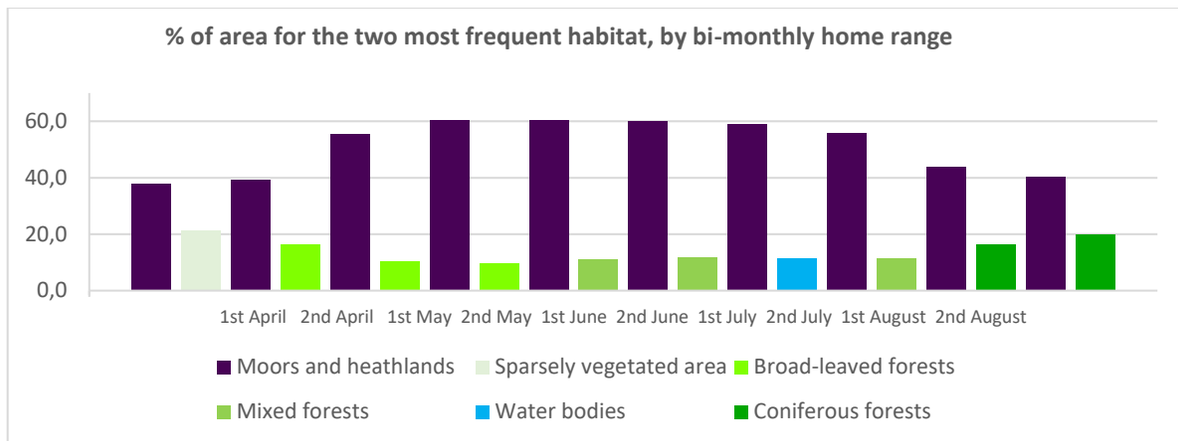
The auto-correlated factors are annual mean temperatures values (1950-1990) and altitudes. This can be explained by the altitudinal gradient, in particular between the high altitudes and low annual temperatures in the north of the studied area, low altitudes and higher temperatures in the south (Hardangervidda plateau). At the scale of the second most extent home range, the correlation is strong and significant ( $r = +0.83$ ;  $p < 0.0001$ ). Then, it is statistically possible to express altitudes distribution as a good proxy for annual temperature distribution. That said, it could be interesting to reprocess the calculation with a more recent dataset (1980-2010 for example) and more accurate temperatures variables (minimal and maximal temperatures).

#### **3.2.2 Ecological habitats and intra-annual (e.g. inter-seasonal) variability of home ranges**

In absolute values of pixels, intermediary seasonal home ranges (early spring, late summer) have the greatest overall area, compared to calving period and full-season home ranges. For example, the most extent home range (1<sup>st</sup> half of April) counts 19736 geostatistical observations (~4480 km<sup>2</sup>) when the smallest one (2<sup>nd</sup> half of June) is about 2262 pixels (~565 km<sup>2</sup>). Indirectly, this fact is shown in the figure 2. Percentages of the two most frequent ecological habitats by fortnight and home ranges during the whole period are here displayed. Colors represent the type of ecological habitat, with a total number of 6. For both April and August fortnights, a relatively balanced distribution is noticed compared to June. Moors and heathland areas are widely represented, with no less than 35% as minima for every home range total areas. Water bodies are represented for the 1<sup>st</sup> July home range (~10%), since any internal differentiation has been made within bi-monthly home ranges.

The  $\chi^2$ -test (performed on the number of pixels of the 9 more represented habitats for the ten fortnights) shows a strong relationship between these variables (observed value: 9873.29, critical value: 92.80,  $p < 0.0001$ ). Some habitats are over-represented during early spring, particularly sparsely vegetated area. During spring and summer, moors and heathland and mixed forests are over-represented. During late summer, coniferous forests and transitional woodlands are over-represented.

Thus, due to the wider extent of reindeer's home ranges during intermediary seasons, habitat diversity appears to be higher when females start to go towards the calving areas. The same can be said during the summer, with mixed forests and later coniferous habitats emerging after the phenological peak of broad-leaved forests followed by the one of moors.



**Fig. 2.** Comparison between reindeer's fortnight home ranges along vegetational season (April to August) and the first two major frequent CLC habitat classes by home range (sources: Corine Land Cover ecological habitat, European Environmental Agency; AdeHabitat package, Callenge, 2006; GRiD-toolbox; credits: R. Courault, 2017)

### 3.2.3 NDVI dissimilarities and selecting specific home ranges

The Kruskal-Wallis test is used to verify if statistical distributions are significantly different. This is the case for the three annual home ranges tested (spring migration: 1<sup>st</sup> half of April; calving area: 2<sup>nd</sup> half of May; fat accumulation area: 2<sup>nd</sup> half of July), totaling 9 distributions. The NDVI values (2007-2008 and 2009) are statistically different ( $p < 0.0001$ ). Between the three years, it is now possible to determine what are the « cold » and « warm » years according to NDVI onset, distribution and central parameter. These chosen years will be studied for the inter-annual home ranges. The specific home range describing migration toward summer pastures (1<sup>st</sup> half of April) under « cold » conditions is 2008, with +0.11 as average of the NDVI. 2009 is the « warm » year with +0.32 as average of NDVI. The home range describing calving period (2d fortnight of May) under « cold » conditions is 2008 (average: +0.45) and 2009 « warm » (+0.56). The specific home range period describing the fat accumulation period under « cold » conditions is 2009 (+0.66), whereas 2007 is « warm » with +0.75.

## 3.3 Inter-annual scale of home ranges: what parameters are related to reindeer estimated densities?

### 3.3.1 Spring as a robust marker for estimated densities of wild reindeers

Table 3 reports Spearman's rho correlations between estimated reindeer's densities (Kernel Utilization Distribution technique) for selected years and relevant explicative factors. Correlations exceeding a threshold of 0.2 ( $p$ -value  $< 0.0001$ ) are shown here. High altitudes are likely associated with high values of reindeer densities during cold years (1<sup>st</sup> fortnight of April and 2<sup>nd</sup> of May, year 2008). High-altitude plant communities, such as bryophytes and terricolous lichens could be preferred, reindeers awaiting seasonal meltdown and vascular plants onset (Klein, 1990). Conversely, « warm » conditions are most likely to correlate with estimated densities of reindeers and NDVI. In that case, correlations describe both negative (Spearman's  $\rho = -0.268$ ,  $p$ -value  $< 0.0001$ ) for April 2009 and positive Spearman's correlation (+0.262,  $p$ -value  $< 0.0001$ ). The negative correlation (1<sup>st</sup> fortnight of April 2008) depicts low densities related to high NDVI values, whereas the positive correlation (2<sup>nd</sup> fortnight of May 2009) depicts high reindeer densities with high NDVI values. Explanations can be found when looking at opportunistic reindeer feeding strategies, in particular

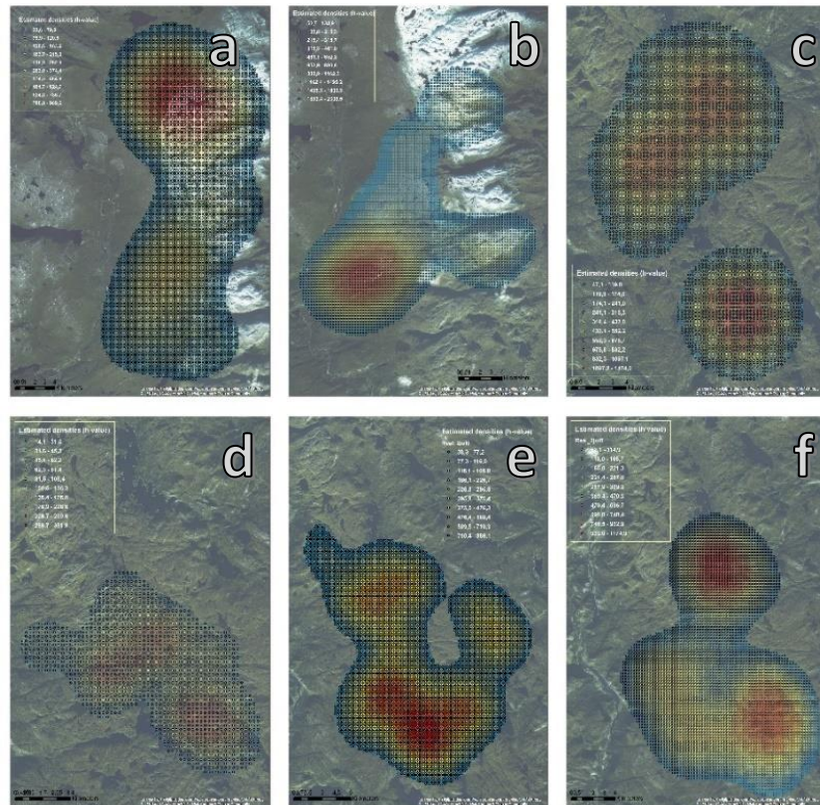
in spring when annual herbaceous just start growing (Skarin et al., 2008). No clear correlation for “fat accumulation” home ranges (2<sup>nd</sup> fortnight of July) has been found. Here, the relative abundance of biomass, shortly after the phenological peak, can explain the lack of relationship.

**Table 3.** Spearman’s rank correlation matrix between densities estimation by annual home ranges and explicative parameters. Only Spearman’s rho > 0.2 and statistical p-value <0.0001 are shown here

Estimation of utilization distribution by annual home ranges/ Explicative factors	Altitudes	NDVI
1st fortnight of April 2008 (cold)	<b>0.374</b>	NA
1st fortnight of April 2009 (warm)	NA	<b>-0.268</b>
2nd fortnight of May 2008 (cold)	<b>0.230</b>	NA
2nd fortnight of May 2009 (warm)	NA	<b>0.262</b>

### 3.3.2 Strong inter-annual variability in estimated densities and phenology between the inter-annual home ranges

In order to explore more thoroughly the relationship between reindeer estimated densities and NDVI values, Kruskal-Wallis tests have been computed. Statistical distributions have been clustered by sub-samples, e.g. ecological habitat classes. Every test calculated for the 9 specific home ranges has rejected the dependence hypothesis between the geostatistical distributions of estimated reindeer densities and NDVI values (9 tests for the 9 specific home ranges). Such variability is displayed in figure 3. The set of maps shows likeness in pattern distribution of estimated densities shapes (ethological/individual behavior, calving area), but also dissimilarities (environmental factors depending on contextual year, spring migration).



**Fig. 3.** Set of maps showing reindeer estimated densities using Kernel Utilization Distribution technique, in a color scale from blue to red. a. 1<sup>st</sup> fortnight of April 2008; b. 1<sup>st</sup> fortnight of April 2009; c. 2<sup>nd</sup> fortnight of May 2008; d. 2<sup>nd</sup> fortnight of May 2009; e. 2<sup>nd</sup> fortnight of July 2007; f. 2<sup>nd</sup> fortnight of July 2009 (Base-map sources: ESRI Dataglobe, Credits: R. Courault, 2017)

## 4. Conclusions

Mounting GRiD toolbox has been incrementally encouraged by the needs of automation processing for the survey of wild reindeer migration and densities in Hardangervidda Plateau (Norway). The wide range spatial behavior of reindeer is an interesting case study to apply this method, more adapted to large territories than simple grid multi-data analysis previously realized by other authors in smaller areas (Jolivet 2014, Bortolamiol et al., 2016). Indeed, by choosing the core point of each grid cell as the common reference to cross all the datasets, the process is realized within a reasonable computation time while avoiding important bias according to the preliminary test realized in our study. Improvements still have to be made to ensure GRiD geometrical reliability, cartographic projection homogenization and minimizing error when extracting large and various raster datasets.

Applying GRiD toolbox for wild reindeer survey during the three growth periods simplified pre-treatments by automatizing grid and creation of mesh of points; and finally datasets fusion into one attribute table. Based on geostatistical observations (pixels) it allowed us to compute several complex statistical analyses, not usually present in GIS software. Based on Open Access datasets, the study firstly shows interesting links between reindeer intra-annual distribution and ecological habitat diversity; secondly relationships between inter-annual variability of estimated densities and abiotic factors, validating thus our second and third hypothesis.

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