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1	The global distribution of leaf chlorophyll content
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3	Croft, H. ^{ab} *Chen, J.M. ^a Wang, R. ^a Mo, G. ^a Luo, S. ^c Luo, X. ^d He, L. ^a Gonsamo, A. ^a Arabian, J. ^{ae}
4	Zhang, Y. ^f Simic-Milas, A. ^g Noland, T.L. ^h He, Y. ⁱ Homolová, L. ^j Malenovský, Z. ^{jk} Yi, Q. ^l Beringer, J. ^m
5	Amiri, R. ⁿ Hutley, L. ^o Arellano, P. ^p Stahl, C. ^q Bonal, D. ^r
6	
7	^a University of Toronto, Department of Geography, Toronto, ON M5S 3G3, Canada
8	*holly.croft@utoronto.ca
9	^b University of Sheffield, Department of Animal and Plant Sciences, Sheffield, S10 2TN, U.K.
10	^c Key Laboratory of Digital Earth Science, Institute of Remote Sensing and Digital Earth, Chinese
11	Academy of Sciences, Beijing 100094, China
12	^d Lawrence Berkeley National Laboratory, Climate and Ecosystem Sciences Division, Berkeley, CA
13	94720, USA
14	^e WWF-Canada, 410 Adelaide Street W, Toronto, ON M5V 1S8, Canada
15	^f Delta State University, Division of Biological and Physical Sciences, Cleveland, MS 38733, USA
16	^g Bowling Green State University, Department of Geology, Bowling Green, OH 43403-0211, USA
17	^h Ontario Ministry of Natural Resources, Ontario Forest Research Institute, 1235 Queen St. E.,
18	Sault Ste. Marie, ON P6A 2E5 Canada
19	ⁱ University of Toronto Mississauga, Department of Geography, 3359 Mississauga Rd,
20	Mississauga, ON L5L 1C6, Canada
21	^j Global Change Research Institute CAS, Bělidla 986/4a, Brno 603 00, Czech Republic

- ²² ^k University of Tasmania, School of Land and Food, Surveying and Spatial Sciences Group, Private
- 23 Bag 76, Hobart, TAS 7001, Australia
- ¹Xinjiang Institute of Ecology and Geography Chinese Academy of Sciences, 818 Beijing South
- 25 Road, Urumqi, Xinjiang 830011, PR China
- ²⁶ ^m The University of Western Australia, School of Earth and Environment (SEE), Crawley WA,
- 27 6009, Australia
- ²⁸ *n* Monash University, School of Earth, Atmosphere and Environment, Clayton VIC, 3800,
- 29 Australia
- ³⁰ ^o Charles Darwin University, School of Environment, Research Institute for the Environment and
- 31 Livelihoods, NT 0909, Australia
- ⁹ Yachay Tech University, School of Geological Sciences and Engineering, Center of Earth
- 33 Observation (CEO), Urcuqui, Ecuador
- ⁹ ⁹ INRA, UMR Ecologie des Forêts de Guyane, Campus Agronomique, BP 709, 97387 Kourou
- 35 Cedex, French Guiana
- ¹ 36 ^r UMR EEF, INRA Université de Lorraine, 54280 Champenoux, France
- 37

38 Abstract

Leaf chlorophyll is central to the exchange of carbon, water and energy between the biosphere and the atmosphere, and to the functioning of terrestrial ecosystems. This paper presents the first spatially continuous view of terrestrial leaf chlorophyll content (Chl_{Leaf}) across a global scale. Weekly maps of Chl_{Leaf} were produced from ENIVSAT MERIS full resolution (300 m) satellite data with a two-stage physically-based radiative transfer modelling approach. Firstly, leaf-level reflectance was derived from top-of-canopy satellite reflectance observations using 4-Scale and SAIL canopy radiative transfer models 45 for woody and non-woody vegetation, respectively. Secondly, the modelled leaf-level reflectance was 46 used in the PROSPECT leaf-level radiative transfer model to derive Chl_{Leaf}. The Chl_{Leaf} retrieval algorithm 47 was validated with measured Chl_{Leaf} data from 248 sample measurements at 28 field locations, and 48 covering six plant functional types (PFTs). Modelled results show strong relationships with field 49 measurements, particularly for deciduous broadleaf forests ($R^2 = 0.67$; RMSE = 9.25 µg cm⁻²; p<0.001), croplands ($R^2 = 0.41$; RMSE = 13.18 µg cm⁻²; p<0.001) and evergreen needleleaf forests ($R^2 = 0.47$; RMSE 50 51 = 10.63 μ g cm⁻²; p<0.001). When the modelled results from all PFTs were considered together, the overall relationship with measured Chl_{Leaf} remained good ($R^2 = 0.47$, RMSE = 10.79 µg cm⁻²; p<0.001). 52 53 This result was an improvement on the relationship between measured Chl_{Leaf} and a commonly used chlorophyll-sensitive spectral vegetation index; the MERIS Terrestrial Chlorophyll Index (MTCI; $R^2 = 0.27$, 54 p < 0.001). The global maps show large temporal and spatial variability in Chl_{leaf}, with evergreen broadleaf 55 forests presenting the highest leaf chlorophyll values with global annual median of 54.4 μ g cm⁻². Distinct 56 57 seasonal Chl_{Leaf} phenologies are also visible, particularly in deciduous plant forms, associated with 58 budburst and crop growth, and leaf senescence. It is anticipated that this global Chl_{Leaf} product will make 59 an important step towards the explicit consideration of leaf-level biochemistry in terrestrial water, 60 energy and carbon cycle modelling.

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Key words: Radiative transfer, 4-Scale, SAIL, PROSPECT, leaf biochemistry, MERIS, satellite, remote
sensing, leaf physiology, carbon cycle, ecosystem modelling, phenology

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69 **1.0** Introduction

70 Chlorophyll molecules facilitate the conversion of absorbed solar radiation into stored chemical energy, 71 and the exchange of matter and energy fluxes between the biosphere and the atmosphere. Our ability 72 to accurately model these fluxes is important to forecasting carbon dynamics, within the context of a 73 changing climate. However, within conventional carbon modelling approaches, the parameterisation of 74 vegetation structure and physiological function over both spatial and temporal domains, with an 75 acceptable level of accuracy remains challenging (Groenendijk et al. 2011, Houborg et al. 2015). Within 76 such modelling approaches, leaf area index (LAI) is a core biophysical parameter used to represent 77 vegetation density, seasonal phenology and the fraction of absorbed PAR by vegetation that is 78 converted to biomass (Bonan et al. 2011). The ecological importance of LAI has led to well-validated 79 datasets of LAI maps at global scales and fine spatial resolution (~1 km) (Baret et al. 2013, Deng et al. 80 2006). However, recent studies have found that a temporal decoupling between vegetation structure 81 and function can occur (Croft, Chen, and Zhang 2014b, Croft, Chen, Froelich, et al. 2015, Walther et al. 82 2016), particularly in deciduous vegetation with a strong seasonal phenology. Chlorophyll molecules 83 comprise an important part of a plant's "photosynthetic apparatus" (Peng et al. 2011), through the 84 harvesting of light and the production of biochemical energy for use within the Calvin-Benson cycle 85 (Porcar-Castell et al. 2014). Leaf chlorophyll content (Chl_{Leaf}) therefore represents a plant's physiological 86 status, and is closely related to plant photosynthetic function; demonstrating a strong relationship to 87 plant photosynthetic capacity (Vcmax) (Croft et al. 2017). Neglecting to consider chlorophyll phenology 88 within carbon models can lead to an overestimation of the amount of plant carbon uptake at the start 89 and end of a growing season in deciduous vegetation (Croft, Chen, Froelich, et al. 2015, Luo et al. 2018). 90 The incorporation of inter- and intra- annual variations of chlorophyll within ecosystem models has been 91 shown to improve the simulations of carbon and water fluxes (Luo et al. 2018, He et al. 2017).

93 Global efforts to map Chl_{Leaf} have been hampered by the complexity in the relationship between 94 satellite-derived canopy reflectance and plant biophysical and biochemical variables. Thus far, satellite 95 remote sensing applications to map Chl_{Leaf} have largely been limited to empirical methods, via the 96 derivation of statistical relationships between spectral vegetation indices (VIs) and leaf or canopy 97 chlorophyll content (Le Maire, François, and Dufrêne 2004, Sims and Gamon 2002). Indices that include 98 'red-edge' wavelengths (690–740 nm) are the most strongly related to Chl_{Leaf} (Croft, Chen, and Zhang 99 2014a, Malenovský et al. 2013), due to the ready saturation of chlorophyll absorption bands when Chl_{Leaf} 100 exceeds \sim 30 µg cm⁻² (Croft and Chen 2018). Some studies have shown promising results using empirical 101 methods (Datt 1998, Haboudane et al. 2002). However, this is usually achieved at local scales, within 102 closely related species (Blackburn 1998) or for uniform, closed canopies, where the vegetation stand 103 essentially behaves as a 'big leaf' (Gamon et al. 2010), and where contributions from other variables, 104 such as background vegetation and non-photosynthetic elements are low. At the leaf level, variations in 105 internal leaf structure, leaf thickness and water content, differentially affect leaf reflectance (Serrano 106 2008, Croft, Chen, and Zhang 2014a). At the canopy scale, vegetation architecture including LAI, foliage 107 clumping, stand density, non photosynthetic elements and understory vegetation, in addition to the 108 sun-view geometry, affect measured reflectance factors (Demarez and Gastellu-Etchegorry 2000, 109 Verrelst et al. 2010, Malenovský et al. 2008).

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An alternative satellite-based approach for deriving Chl_{Leaf} from top-of-canopy reflectance data is through the use of physically-based radiative transfer models. Radiative transfer models provide a direct physical relationship between canopy reflectance and Chl_{Leaf} because they are underpinned by physical laws that determine the interaction between solar radiation and the vegetation canopy. Leaf-level estimation of foliar chlorophyll is achieved by coupling a canopy model and leaf optical model, in a twostep process (Croft and Chen 2018): firstly to derive leaf level reflectance from canopy reflectance and

then to derive leaf pigment content from the modeled leaf reflectance (Zhang et al. 2008, Croft et al. 117 118 2013, Zarco-Tejada et al. 2004). A number of canopy models have been used for this purpose, ranging 119 from turbid medium models (e.g. SAIL) (Verhoef 1984), hybrid geometric optical and radiative transfer 120 models (e.g. 4-SCALE (Chen and Leblanc 1997), GeoSAIL (Huemmrich 2001), DART (Gastellu-Etchegorry, 121 Martin, and Gascon 2004)) in which the turbid media are constrained into a geometric form (i.e. a leaf, 122 shoot, branch and/or crown), and ray-tracing techniques (FLIGHT) (North 1996). At the leaf level, the 123 most widely used leaf optical model is PROSPECT (Jacquemoud and Baret 1990), which has been 124 extensively applied and validated across a wide range of plant species (Malenovský et al. 2006), due to 125 the small number of input parameters required and its ease of inversion. Previous research has 126 demonstrated the strength of this physically-based method for a number of different ecosystems (Zarco-127 Tejada et al. 2004, Zhang et al. 2008, Croft et al. 2013, Demarez and Gastellu-Etchegorry 2000, Houborg 128 and Boegh 2008). However, this approach has yet to be applied at the global scale.

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130 This paper presents the first global Chl_{Leaf} map from satellite data using physically-based radiative 131 transfer models. Chl_{Leaf} is defined on a leaf-area basis, as chlorophyll content per half the total surface 132 leaf area. Expressing Chl_{Leaf} by leaf area (as opposed to by dry mass) is the closer representation of what 133 is directly measured by a satellite instrument, and is most appropriate for linking Chl_{Leaf} to ecosystem 134 processes, such as carbon and water fluxes in relation to surfaces (Wright et al. 2004). Chl_{Leaf} is modelled 135 from ENVISAT MEdium Resolution Imaging Spectrometer (MERIS) 300 m reflectance data in a two-step 136 modelling approach, using coupled canopy and leaf radiative transfer models. Modelled Chl_{Leaf} results 137 were subsequently validated using measured ground data at a range of different field sites over six 138 different plant functional types (PFTs). Chl_{Leaf} maps are produced at the global scale every seven days for 139 an entire calendar year 2011 in order to provide spatially- and temporally-distributed leaf chlorophyll 140 content, and consequent physiological information for ecosystem modelling and ecological applications.

142 **2.0 Methods**

143 **2.1 Ground data and validation sites**

We used measured leaf chlorophyll data from 248 sampling measurements within 28 field locations covering six PFTs (Figure 1 and Table 1) to validate the Chl_{Leaf} retrieval algorithm. These data included 49 measurements in deciduous broadleaf forests (DBF), 9 measurements in evergreen broadleaf forests (EBF), 100 measurements in needleleaf forests (ENF), 55 measurements in croplands (CRP), 28 measurements in grassland (GRS) and 7 measurements in shrublands (SHR).



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Figure 1: Locations of the twenty-eight field locations used for leaf chlorophyll content validation.

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For each field location, the number of discrete sites or dates over which individual ground measurements were taken are reported in Table 1. The number of replicates and spatial sampling varied between the field sites, due to the nature of data collection in individual projects and between researchers, and represents the large number of sources that have come together to produce this work. Table 1: Details of the ground measurements of leaf chlorophyll content used for validation used in this study, including the sampling location (Lat = latitude, Long = longitude) and sampling year, dominant species, plant functional type (PFT), mean site leaf area index (LAI) value and chlorophyll determination

160 *method (Chl. method).*

Site name	Country	Sampling years	Lat/ Long*	PFT	Dominant species	Mean LAI	Mean Chl. (μg cm ⁻²)	No. of sites/ dates	Chl. method†	Reference
Haliburton	Canada	2004	45.24 -78.54	DBF	Sugar maple	5.5	38.3	8	Lab	Zhang et al. (2007)
Borden Forest	Canada	2013-15	44.32 -79.93	DBF	Red maple	4.2	37.8	13	Lab	Croft et al. (2015)
Bioindicators DBF	Canada	2002	46.84, -81.41	DBF	White birch, poplar	-	34.3	26	Lab	-
Sudbury DBF	Canada	2007	47.16, -81.71	DBF	Trembling aspen	4.3	48.8	2	Lab	Simic et al. (2011)
Amazon FG	French Guyana	2007-08	5.28, -52.92	EBF	-	6.7	62.1	3	Chl. meter	Rowland et al. (2014)
Amazon Ecuador	Ecuador	2012	-0.18, -76.36	EBF	-	-	53.6	6	Field spec.	Arellano et al. (2017)
Sudbury ENF	Canada	2007	47.18, -81.74	ENF	Black spruce		29.0	10	Lab	Simic et al. (2011)
Chapleau	Canada	2012	47.58, -83.01	ENF	Jack pine	3.3	49.3	8	Lab	Croft et al. (2014)
Bioindicators ENF	Canada	2001-02	47.11, -81.82	ENF	Black spruce, Jack pine	-	32.8	60	Lab	-
Sudbury Zhang	Canada	2003-04	47.16, -81.74	ENF	Black spruce	3.2	30.4	18	Lab	Zhang et al. (2008)
Le Casset	France	2008	44.98 <i>,</i> 6.48	ENF	European larch	-	60.8	2	Lab	Homolová et al., (2013)
Bily Kriz	Czech Republic	2004 <i>,</i> 2006	49.50, 18.54	ENF	Norway spruce	8.2	39.8	2	Lab	Homolová et al. (2013); Malenovský et al. (2013)
Col du Lautaret Terrasses	France	2008	45.04 <i>,</i> 6.35	GRS	Mixed alpine sp.	-	38.5	3	Lab	Homolová et al. (2014)
Roche Noir	France	2008	45.06, 6.38	GRS	Mixed alpine sp.	-	39.3	5	Lab	Pottier et al. (2014)
GRS National Park	Canada	2012-14	49.16, -107.56	GRS	Mixed-grass prairie	1.0	35.4	20	Lab	Tong and He (2017)
Stratford Wheat	Canada	2013	43.45, -80.86	CRP	Wheat	2.2	38.7	5	Lab	Dong et al. (2017)

	_		43.46,			2.7				
Stratford Corn	Canada	2013	-80.81	CRP	Maize	2.7	42.4	10	Lab	Dong et al. (2017)
Mosuowanzhen	China	2011	44.19 <i>,</i> 85.49	CRP	Cotton	1.87	64.3	15	Lab	Yi et al., (2014)
Sele River Plain	Italy	2009	40.52 <i>,</i> 15.00	CRP	Maize, fruit trees	2.3	42.6	29	Chl. Meter •	Vuolo et al. (2012)
Demmin	Germany	2006	53.99, 13.27	CRP	Maize, wheat, Sugarbeet	-	35.4	4	Chl meter	Hajnsek et al. (2006)
Trapani	Sicily	2010	37.64 <i>,</i> 12.85	CRP	Olive trees	1.5	39.6	3	Chl. meter	-
Howard Springs	Australia	2009	-12.49, 131.15	SHR	Eucalyptus	1.3	57.7	1	Lab	Amiri (2013)
Daly Uncleared	Australia	2009	-14.16, 131.39	SHR	Eucalyptus	0.8	50.1	1	Lab	Amiri (2013)
Daly Regrowth	Australia	2009	-14.13, 131.38	SHR	Eucalyptus	0.9	45.1	1	Lab	Amiri (2013)
Adelaide River	Australia	2009	-13.08, 131.12	SHR	Eucalyptus	0.7	61.5	1	Lab	Amiri (2013)
Dry Creek	Australia	2009	-15.26, 132.37	SHR	Eucalyptus	0.8	45.9	1	Lab	Amiri (2013)
Sturt Plains Shrubland	Australia	2009	-17.15, 133.35	SHR	Eucalyptus	0.0	58.2	1	Lab	Amiri (2013)
Sturt Plains Woodland	Australia	2009	-17.18, 133.35	SHR	Acacia	0.7	45.1	1	Lab	Amiri (2013)

* where multiple sampling sites are present at a given field location, approximate central co-ordinates are given.
DBF = Deciduous broadleaf, ENF = Evergreen needleleaf, EBF = Evergreen broadleaf, GRS = Grassland, CRP =
Cropland, SHR = Shrubland. †Lab = laboratory extraction; Chl meter = handheld optical meter; Field spec = field
spectrometer. • indicates species specific chl. meter calibration equations.

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Methods of chlorophyll measurement presented in Table 1 were either through laboratory analysis techniques (Lab) or via optical methods (Croft and Chen 2018), i.e. a handheld chlorophyll meter (Chl. meter) or a field spectroradiometer (Field spec.). Where species specific calibration equations were used, further details of the calibration equations can be found in the corresponding referenced paper. The field spectrometer-based retrieval (Arrellano et al., 2017) used the PROSPECT radiative transfer model (Jacquemoud et al., 1990) to estimate chlorophyll content. It is recognised that the different 172 methods of chlorophyll determination may lead to a certain degree of variability between validation 173 measurements. The algorithm was validated according to the closest available ENVISAT MERIS satellite 174 date and location to the field sampling date. Due to the relative scarcity of available in situ Chl_{Leaf} data, 175 the existing validation data were maximised by using data collected in years shortly preceding or 176 following the MERIS operational time frame (i.e. earlier than 2002 or later than 2011; Table 1). In this 177 case, the correct day of year (DOY) from the closest calendar year within the 2002-2011 time period, 178 was used. The validation results (Section 3) are partitioned to indicate whether the results are from the 179 current year or closest matched year.

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181 2.2 Satellite data

182 2.2.1 MEdium Resolution Imaging Spectrometer (MERIS) satellite data

183 ENIVSAT MERIS satellite-derived reflectance data was selected as the input remote sensing product for 184 this study, due to the sampling of chlorophyll-sensitive red-edge bands, a short temporal revisit time 185 (every 2-3 days) for global application, medium spatial resolution (300 m) and its high radiometric 186 accuracy (Curran and Steele 2005). We used the full resolution (FR) surface reflectance (SR) product, 187 which is produced as a 7-day temporal synthesis from data collected at the original 2-3 day revisit 188 frequency. The global SR time series are produced by a series of pre-processing steps within the MERIS 189 pre-processing chain, including radiometric, geometric, bidirectional reflectance distribution function 190 (BRDF), pixel identification, and atmospheric correction with aerosol retrieval. The 7-day product normalises reflectance to nadir view, and the solar zenith angle is that of 10h00 local time for the 191 192 median day of the compositing period. MERIS reflectance data is provided in 13 bands (spectral 193 resolution = ~10 nm) in the visible, red-edge and near infra-red bands, with the atmospheric bands 194 (bands 11 and 15) removed. The MERIS FR surface reflectance global time-series covers the 2003-2012 195 time period.

197 2.2.2 Landcover map

A global land cover map produced from 300 m spatial resolution MERIS data, as part of the European Space Agency Climate Change Initiative (CCI-LC) project, from the 5-year period (2008-2012) was used to define PFTs. The legend is based on the UN Land Cover Classification System (LCCS) with the view to be compatible with the Global Landcover 2000 (GLC2000), GlobCover 2005 and 2009 products.

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203 2.2.3 Leaf area index (LAI)

204 Copernicus Global Land Service GEOV1 LAI product derived from SPOT-VGT satellite (Baret et al., 2013) 205 was used in the algorithm, with a global LAI coverage from 1999 to the present at 10-day temporal 206 intervals and a spatial resolution of 1 km. The GEOV1 LAI product is derived from the CYCLOPES v3.1 and 207 MODIS c5 biophysical products, through a neural-network machine-learning algorithm (Baret et al. 208 2013). The selection of CYCLOPES v3.1 and MODIS c5 products takes advantage of previous 209 development efforts and capitalises on the performances of each product (Camacho et al. 2013). As part 210 of the processing chain, the LAI product is reprojected onto the Plate-Carrée 1/112° grid, temporally 211 smoothed, interpolated at the 10-day frequency, and re-scaled to fit the expected range of variation 212 (Verger et al. 2015). GEOV1 LAI products consider clumping as a weighted contribution of CYCLOPES and 213 MODIS products, where MODIS LAI accounts for clumping at plant and canopy scales (Knyazikhin et al. 214 1998), with the exception of needleleaf forests, for which shoot clumping is not accounted for. In the 215 CYCLOPES algorithm, landscape clumping is represented by considering fractions of mixed pixels (Baret 216 et al. 2013). Recent validation studies indicated that GEOV1 outperformed most existing products in 217 both accuracy and precision (Camacho et al. 2013). Validation results showed good spatial and temporal 218 consistency and accuracy, smooth and stable temporal profiles, good dynamic range with reliable 219 magnitudes for bare areas and dense forests (Camacho et al. 2013). The GEOV1 LAI product was

selected for this project, because it met the following criteria: a high temporal frequency (every 10 days), acceptable spatial resolution (1 km), a long archive of data covering the complete MERIS mission (2002-2012), and strong product validation.

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224 2.3 Algorithm development

225 To derive leaf chlorophyll content from remote sensing data at the global scale, a two-step modelling 226 approach was selected. The first step in the algorithm is the modelling of leaf-level reflectance spectra 227 from satellite-derived canopy reflectance data, using physically-based radiative transfer models to account for the influence of canopy architecture, image acquisition conditions and background 228 229 vegetation on canopy reflectance. The second step was to retrieve leaf chlorophyll content from the 230 modelled leaf reflectance derived in Step 1, using a leaf optical model. A two-step inversion method is 231 favoured over a coupled one-step inversion because the output of each stage can be assessed 232 individually, and may be validated against measured leaf-level reflectance data at field sites (Croft et al. 233 2013, Zhang et al. 2008). This physically-based canopy inversion method has been previously 234 successfully demonstrated using different combinations of canopy and leaf models (Croft et al. 2013, 235 Moorthy, Miller, and Noland 2008, Zarco-Tejada et al. 2004, Kempeneers et al. 2008).

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For the first step, two canopy reflectance models were selected, according to the structural characteristics of the vegetation present. For spatially heterogeneous 'clumped' vegetation types (i.e. deciduous and coniferous trees, shrubs), the 4-Scale geometrical–optical model (Chen and Leblanc 1997) was used (see Section 2.3.1). For homogenous canopies that can be treated as one-dimensional (1D) turbid media, such as agricultural crops, we used the SAIL model (Verhoef 1984) (Section 2.3.2). To invert the canopy radiative transfer models, individual LUTs for different PFTs were created, based on variable and fixed input parameters (Table 2 and Table 3). The LUT approach was selected to optimise

244 computational resources and reduce problems associated with appearances of local minima, given 245 sufficient sampling of the variable space (Jacquemoud et al. 2009). Whilst these structural 246 parameterisations are important, their influence on canopy reflectance is mediated by LAI, which is the 247 dominant driver of modelled canopy reflectance (Zhang et al. 2008 and Section 2.4). The availability of 248 accurate and well validated global-scale LAI products facilitated the treatment of LAI as a variable 249 parameter, which was derived from the GEOV1 LAI product (Section 2.2.3) at 10 day intervals and 1 km 250 spatial resolution. In the second step, leaf chlorophyll content was retrieved using a leaf optical model 251 (PROSPECT; Jacquemoud and Baret 1990) (Section 2.3.3) from the modelled leaf reflectance that was 252 derived in step 1.

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254 **2.3.1** Step 1: Leaf reflectance inversion - 4-Scale model

255 The 4-Scale model (Chen and Leblanc 1997) was selected for the inversion of leaf level reflectance from 256 satellite-derived images in forested and spatially clumped ecosystems. 4-Scale considers the three-257 dimensional (3D) spatial distribution of vegetation groups and vegetation crown geometry, in addition 258 to the structural effects of branches and scattering elements. In closed canopies, structural variables 259 such as crown shape and clumping of foliage elements are dominant, whereas in open canopies the 260 effects of background reflectance and shadows prevail. The 4-Scale model simulates the Bidirectional 261 Reflectance Distribution Function (BRDF) based on canopy architecture described at four scales: 1) 262 vegetation grouping, 2) crown geometry, 3) branches, and 4) foliage elements (Chen and Leblanc 2001). 263 A crown is, therefore, represented as a complex medium, where mutual scattering occurs between 264 shoots or leaves. Consequently, sunlit foliage can be viewed on the shaded side and shadowed foliage 265 on the sunlit side. Reflected radiance from different scene components is calculated by first separating 266 sunlit and shaded components through first-order scattering, and then adding multiple scattering from 267 subsequent interactions with vegetation elements or background material (Chen and Leblanc 2001). To

268	model canopy reflectance, the 4-Scale model was run in the forward mode, with variable (LAI, solar
269	zenith angle and view zenith angle) and fixed structural parameters and background reflectance spectra
270	(Table 2), set according to ground measurements and values reported in the literature (Chen and
271	Leblanc 1997, Leblanc et al. 1999). The element clumping index (ΩE) represents vegetation clumping at
272	scales larger than the shoot, and is associated with canopy architecture and structural variables, such as
273	crown size and vegetation density. ΩE is an important parameter for estimating radiation interception
274	and distribution in plant canopies (Chen, Menges, and Leblanc 2005). The non-random spatial
275	distribution of trees is simulated using the Neyman type A distribution to permit clumping and
276	patchiness within a forest stand (Chen and Leblanc 1997).
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292 Table 2: Fixed and variable parameters used in the 4-Scale model for LUT generation.

	Broadleaf	Needleleaf	Chuuhland	References	
	trees	trees	Shrubland		
LAI (m ² m ^{- 2})	0.1-8	0.1-8	0.1-8	Baret et al., (2013)	
Solar zenith angle (°)	20-70	20-70	20-70	MERIS metadata	
View zenith angle (°)	0	0	0	MERIS metadata	
Relative azimuth angle (°)	0	0	0	MERIS metadata	
Stick height (m)	10	10	3	Los et al. (2012)	
Crown height (m)	8	10	7	Los et al. (2012)	
Crown shape	Spheroid	Cone & cylinder	Spheroid	Chen and Leblanc (1997)	
Tree density (trees/ha)	1400	3000	1000	Chen et al. (1997) & Leblanc et al. (1999)	
Crown radius (m)	1.25	1.0	1.25	Chopping et al. (2008), Evans et al. (2015) & Thorpe et al. (2010)	
Neyman grouping	3	4	3	Leblanc et al. (1999)	
Clumping index (ΩE)	0.90	0.80	0.80	Pisek et al. (2011)	
Needle to shoot ratio (yE)	1	1.4	1	Pisek et al. (2011)	
Foliage element width (m)	0.15	0.1	0.15	Chen and Cihlar (1995) & Leblanc et al. (1999)	
Background composition	Green vegetation and soil	Green vegetation and soil	Dry grasses and soil	Observations/ measurements	

293 Broadleaf/needleleaf classes include respective deciduous and evergreen plant functional types.

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According to the specified input parameters, the 4-Scale model calculates canopy reflectance (*ρ*) as a
 linear summation of four components:

297

$$\rho = \rho_{PT\lambda}F_{PT} + \rho_{ZT\lambda}F_{ZT} + \rho_{PG\lambda}F_{PG} + \rho_{ZG\lambda}F_{ZG}$$
 [Eq. 1]

299

300 where the reflectance factors from each scene component are: sunlit vegetation ($\rho_{PT\lambda}$), shaded 301 vegetation ($\rho_{ZT\lambda}$), sunlit ground ($\rho_{PG\lambda}$) and shaded ground ($\rho_{ZG\lambda}$), and F_{PT} , F_{PG} , F_{PG} and F_{ZT} represent the 302 probability of viewing each component, respectively (Chen and Leblanc 2001). The 4-Scale model includes a multiple scattering scheme, for 2nd order scattering and above, through computing the interactions among the four scene components based on view factors from one component to the other (Chen and LeBlanc, 2001). In this unique way, the geometrical effects propagate to all orders of multiple scattering. In order to derive leaf reflectance ($\rho_{L\lambda}$) from sunlit crown reflectance (ρ_{PT}), the enhancement of both sunlit and shaded reflectance due to multiple scattering is accounted for using a multiple scattering factor (Zhang et al. 2008, M factor; Croft et al. 2013, Simic, Chen, and Noland 2011). The M factor can be calculated using the output sunlit and shaded scene components from 4-Scale:

310

311
$$M_{\lambda} = \frac{\rho_{\lambda} - \rho_{PG_{\lambda}}F_{PG}}{\rho_{L_{\lambda}}F_{PT}}$$
 [Eq. 2]

312

Finally, the M factor (Eq. 2) is used to derive leaf reflectance from canopy-level satellite reflectance ($\rho_{satellite}$), by converting sunlit crown reflectance into sunlit leaf reflectance, and allowing the inclusion of the less variable shaded components (Eq. 3).

316

317
$$\rho_{L\lambda} = \frac{\rho_{satellite\lambda} - \rho_{PG\lambda}F_{PG}}{M_{\lambda}F_{PT}}$$
 [Eq. 3]

318

319

320 **2.3.2** Step 1: Leaf reflectance inversion - SAIL model

The SAIL radiative transfer model (Verhoef 1984) was used to derive leaf reflectance from satellitederived images in cropland and grassland ecosystems, where the distribution of foliage approaches randomness, and information on canopy structural variables such as crown shape and the clumping of foliage elements is not applicable. Turbid medium models such as SAIL (Verhoef, 1984) assume that the canopy is composed of homogeneous, horizontal layers of Lambertian scatterers randomly distributed in space. The SAIL model is one of the most popular and well validated models used in agricultural systems (Jacquemoud et al. 2009, Darvishzadeh et al. 2008, Clevers and Kooistra 2012). SAIL is based on Suit's model (Suits 1971), which is founded on a set of four differential equations: (1) diffuse incoming flux, (2) diffuse outgoing flux, (3) direct solar flux, and (4) flux with radiance in the direction of remote sensing observation. Table 3 details the fixed and variable parameters used within the SAIL model, based on field observations and literature (Privette et al. 1996, Darvishzadeh et al. 2008) that were used to model canopy reflectance for the LUT-based inversion.

333

	Cropland	Grassland	References	
Leaf area index	0.1 - 8	0.1 - 8	Baret et al., (2013)	
Solar zenith angle (°)	20-70	20-70	MERIS metadata	
View zenith angle (°)	0	0	MERIS metadata	
Relative azimuth angle (°)	0	0	MERIS metadata	
Leaf inclination distribution function	Ellipsoidal	Erectophile	Clevers and Kooistra (2012) & Zou and Mõttus (2015)	
Hot spot parameter (m m ⁻¹)	0.1	0.1	Darvishzadeh et al. (2008) & Vohland and Jarmer (2008)	
Soil factor (dimensionless)	1	0.4	Observations/ measurements	

334 Table 3: Fixed and variable parameters used in the SAIL model for LUT generation.

335

336

The leaf inclination distribution function (LIDF) describes the frequency distribution of leaf inclination angles in different directions. For agricultural crops, LIDF is ellipsoidal (Campbell, 1986), with the mean leaf inclination angle set at 40°, in line with values measured within previous studies (Zou and Mõttus 2015, Liu, Pattey, and Jégo 2012). For grasslands the LIDF is erectophile (LiDFa = -1, LiDFb = 0) (Clevers and Kooistra 2012, Sandmeier et al. 1999, He, Guo, and Wilmshurst 2007, Verma et al. 2017, Migliavacca et al. 2017). The hot spot (Hs) parameter is defined as the ratio between the average leaf size and the height of the canopy (Darvishzadeh et al. 2008, Verhoef 1985). The soil factor (Sf) accounts for variations in the brightness of the soil background, where a value of 0 denotes a wet soil and 1 a dry soil. A LUT was generated from the fixed and variable parameters shown in Table 3, which contains the ratio of the modelled canopy reflectance from the SAIL model to the SAIL input leaf reflectance as a scaling factor (S Factor_{λ}). Modelled leaf reflectance ($\rho_{L\lambda}$) is retrieved from satellite canopy reflectance ($\rho_{satellite\lambda}$) according to Equation 4.

349

350

$$\rho_{L_{\lambda}} = \rho_{Satellite_{\lambda}} * S Factor_{\lambda}$$
 [Eq. 4]

351

352 2.3.3 Step 2: Deriving leaf chlorophyll content – PROSPECT model

353 The leaf radiative transfer model PROSPECT (Jacquemoud and Baret 1990, Feret et al. 2008) is used to derive Chl_{Leaf} from the modelled leaf reflectance spectra generated in step one (Section 2.3.1 and 2.3.2). 354 355 In PROSPECT-5, leaf reflectance and transmittance (400-2500 nm) are defined as a function of six 356 parameters: structure parameter (N), chlorophyll (a+b) concentration (C_{ab}), carotenoid content (Car), 357 brown pigment (C_b), dry matter (C_m) and equivalent water thickness (C_w). Absorption is calculated as the 358 linear summation of the specific absorption coefficients of the biochemical constituents and their 359 respective concentrations. PROSPECT has had widespread validation across a large number of 360 vegetation species and plant functional types (Croft, Chen, Zhang, et al. 2015, Demarez and Gastellu-361 Etchegorry 2000, Darvishzadeh et al. 2008, Malenovský et al. 2006). Whilst previous studies inverted 362 PROSPECT by iteratively minimising a merit function (Croft et al., 2013), the large volume of data used in 363 this study precluded this option. In order to optimise computational resources, using a vegetation index 364 (VI) based inversion approach. PROSPECT was run in the forward mode to derive chlorophyll-sensitive 365 spectral vegetation indices, according to incrementing chlorophyll content values. The leaf biophysical 366 parameters were varied for each PFT (Table 4), based on measured values and within reported ranges in 367 the literature.

369

Table 4: Leaf parameters used in the PROSPECT model, along with the vegetation index (VI) derived from forward-modelled reflectance spectra for each plant functional type. C_{ab} = chlorophyll content; N = structural parameter; Car = carotenoid content; C_b = brown matter; C_m = dry matter content; C_w = equivalent water thickness.

	С _{аь} (µg cm ⁻²)	N (unitless)	Car (µg cm ⁻²)	Cь	Cm (g cm ⁻²)	C _w (g cm ⁻²)	Vegetation index	Reference
DBF	0-100	1.2	Chl/7	0	0.005	0.01	MERIS _{SR}	Féret et al. (2011)
EBF	0-100	1.8	Chl/7	0	0.005	0.01	MERISsr	Arellano et al. (2017)
DNF	0-100	2.8	Chl/7	0	0.05	0.01	MERIS _{ND}	De Santis et al. (2009) & Kötz et al. (2004)
ENF	0-100	2.8	Chl/7	0	0.05	0.01	MERIS _{ND}	De Santis et al. (2009) & Kötz et al. (2004)
SHR	0-100	1.8	Chl/7	0	0.005	0.01	MERIS _{ND}	Enrique et al. (2016) & Sow et al. (2013)
CRP	0-100	1.4	Chl/7	0	0.015	0.01	MERIS _{ND}	Jacquemoud et al. (2000)
GRS	0-100	1.2	Chl/7	0	0.005	0.01	MERIS _{ND}	Darvishzadeh et al. (2008)

375

The VIs used in the VI-based PROSPECT inversion are focussed on the red-edge reflectance bands that are resistant to saturation at high chlorophyll values. Croft et al., (2014a) found strong relationships between normalised difference (ND) and simple ratio (SR) vegetation indices with chlorophyll content (Zarco-Tejada et al. 2001, Mutanga and Skidmore 2004), which were modified to MERIS red-edge spectral bands as follows:

381

382

$$MERIS_{ND} = \frac{R_{754} - R_{709}}{R_{754} + R_{709}}$$
[Eq. 5]

$$MERIS_{SR} = \frac{R_{754}}{R_{709}}$$
 [Eq. 6]

where *R* is reflectance at given wavelength λ in nm (*R* λ). Regression equations were then generated between incrementing chlorophyll content and the modelled VI from PROSPECT (Figure 2), which are then applied to the modelled leaf level reflectance derived from canopy model inversion in Step 1 (Sections 2.2.1 and 2.2.2) to derive the final Chl_{Leaf} values.



Figure 2: Regression equations for the PROSPECT-based leaf chlorophyll inversion. The shaded grey area represents the potential uncertainty of the forward modelled vegetation index values, according to a range of measured PROSPECT parameter values from published data. For all plant functional types apart from needleleaf forests, the range in input values is two standard deviations either side of the mean from the LOPEX dataset. For needleleaf forests uncertainty values from Kötz et al. (2004) were used.

399 The shaded grey area in Figure 2 indicates the degree of uncertainty arising from the VI-inversion step, 400 according to the fixed biophysical parameters that are used to parameterise the PROSPECT model. 401 Measured N, C_m and C_w values from the LOPEX published dataset of leaf reflectance and biophysical 402 parameters (Hosgood et al. 1995) were used for DBF, SHR, GRS, CRP species. For the needleleaf and EBF 403 species, for which large measured datasets are not so widely available, we used parameter uncertainty 404 values from Kötz et al. (2004) and standard deviations from Ferreira et al. (2013) for needleleaf and EBF 405 species, respectively. For DBF, SHR, GRS, CRP and EBF species, the PROSPECT model was run in the 406 forward mode for input parameter values that represented two standard deviations of the mean N, Cm 407 and C_w values within the published datasets (represented by grey shading), for each PFT individually. The 408 orange line represents the relationship between the VI and Chl_{Leaf} using the structural parameters in 409 Table 4. The regression equation shown is the empirical model that is used to convert modelled leaf 410 level reflectance derived in Step 1 (Sections 2.3.1 and 2.3.2) to Chl_{Leaf}.

411

412 The VI-based PROSPECT inversion method was assessed against the conventional PROSPECT inversion 413 method of iteratively minimising a merit function (Feret et al., 2008), for the ground validation sites 414 detailed in Table 1. The relationship between the modelled Chl_{Leaf} from the VI-inversion and modelled Chl_{Leaf} from the merit function inversion was R² = 0.74 (percentage bias = 20% and RMSE = 11.01 µg cm⁻ 415 416 ²). Moreover, the VI-inversion increased the relationship between modelled and measured Chl_{Leaf} by 20% 417 (p<0.001), when compared to the merit function Chl_{Leaf} inversion. Using VIs centred on the chlorophyll-418 sensitive red-edge rather than inverting the model across the full leaf spectra reduces the confounding 419 influence of other biophysical variables, such as leaf structure and carotenoid content. Several studies 420 have found a high redundancy of wavelength channels in vegetation studies (Jacquemoud et al., 1995; 421 Simic and Chen, 2008, Croft et al. 2015). Thenkabail et al. (2004) reported that the data volume can be

reduced by 97% when hyperspectral wavebands are reduced to the first five principal components, and
still explain close to 95% of data variability. A reduction in spectral data may therefore permit a better
model fit over the more sensitive and relevant wavelength ranges.

- 425
- 426

2.4 Canopy model inversion sensitivity analysis

427 A sensitivity analysis was undertaken to assess the impact of canopy structural parameters values 428 differing from the fixed value used in the inversion algorithm on the modelled Chl_{Leaf} value. Four of the 429 major PFTs with different structural parameterisations and that employed radiative transfer models 430 (DBF, ENF, CRP, GRS) were tested. All values were held constant according to the values used in the 431 chlorophyll inversion algorithm (Tables 2 and 3), with one parameter incrementally perturbed. The value 432 zero represents the condition under which the structural parameters are consistent with the values used 433 in the main chlorophyll inversion algorithm, as outlined in Tables 1 and 2 (i.e. the reference case). In the 434 case of LAI, which is a variable parameter in the algorithm we used a representative value (LAI = 2.5 435 (GRS), 4.0 (CRP), 5.0 (ENF), 4.0 (DBF)) as found in a global analysis by Asner et al., (2003), for each PFT to 436 avoid bias from an extreme case. Solar and view zenith angles were held constant at 40° and 0°, 437 respectively. To achieve comparability between structural parameters, results are scaled to a 438 percentage difference from the reference case. The chlorophyll values are shown as an absolute 439 difference from the reference case, in order to provide an estimate of the impact that deviations in the 440 actual structural parameter values from the one used in the modelling algorithm has on modelled leaf 441 chlorophyll content values. The structural parameters vary slightly between the PFTs and two models 442 (PROSAIL and 4-Scale), according to the structural parameters present in the model.

443



451 LiDFs are imposed on the x axis scale with LIDF names indicated, where Un = Uniform, Ex = Extremophile,

452

Plg = Plagiophile, Pln = Planophile, Sph = Spherical, Er= Erectophile.

454 Figure 3 reveals different sensitivities of modelled Chl_{Leaf} values in response to changing a specific 455 structural parameter. In all cases variations in LAI presented a dominant effect on the modelled 456 chlorophyll content. For crops a 25% difference in LAI from the reference case (i.e. LAI = 3.0 and 5.0), resulted in a change in modelled chlorophyll by -7.8 μ g cm⁻² and +3.76 μ g cm⁻². The asymmetric 457 458 response of the modelled Chl_{Leaf} values to LAI deviations represents the non-linear response of canopy 459 reflectance to increasing LAI, which saturates at higher LAI values. In the GRS results, this saturation 460 does not occur because of the erectophile leaf inclination angle distribution, and because the LAI 461 reference case is lower (LAI = 2.5), resulting in smaller maximum values (GRS + Δ 100% LAI = 5.0, CRP $+\Delta 100\%$ LAI = 8.0). For ENF trees, using the 4-Scale model, variations in canopy density also presented a 462 463 high degree of sensitivity in modelled Chl_{Leaf} values at extreme low canopy density values. A 33% 464 difference in density from the reference case (i.e. 2000 stems/ha and 4000 stems/ha), resulted in a change in modelled chlorophyll by -6.4 μ g cm⁻² and +4.4 μ g cm⁻², while a 66% difference resulted in a -465 466 19.4 μ g cm⁻² and +7.17 μ g cm⁻² change. For ENF and DBF, deviations in other structural parameters (Ω E, 467 crown height and yE for ENF species) presented a smaller impact on the modelled Chl_{Leaf}. A 20% difference in crown height, for example, only affected Chl_{Leaf} values by +1.5 µg cm⁻² and -1.18 µg cm⁻². 468 469 For the SAIL model parameters in CRP plants, in addition to LAI, the leaf angle distribution (fixed value = 40°) also strongly affected modelled Chl_{Leaf} at larger leaf inclination angles, approaching more 470 471 erectophile leaf inclination angles (>60°). From leaf angle distribution = 10° to 50° the imposed 472 variations in modelled chlorophyll were -2.7 μ g cm⁻² to -4.9 μ g cm⁻². This result was also consistent for 473 the GRS results, where the different LIDFs are noted in Figure 3d. The LIDFs exhibited little change until 474 the leaf inclination angle approached those associated with erectophile canopies, with the plagiophile LIDFs exhibiting the largest difference in chlorophyll values from the modelled erectophile reference 475 476 case (+12.3 µg cm⁻²). Changes in the soil factor and the hotspot parameter induced very little differences 477 in modelled Chl_{Leaf}. For example, in CRP plants a +100% change in the hotspot value (Hs = 0.2) only 478 resulted in a -1.2 μg cm⁻² in Chl_{Leaf}, confirming the findings of Vohland and Jarmer (2008). Deviations in 479 Chl_{Leaf} values that are close to zero, with increasing percentage change in the structural parameter 480 indicates a greater tolerance of the model algorithm to differences in the actual structural parameter 481 value at a given site, to the value that is fixed in the model. The inclusion of LAI as a variable parameter 482 mitigates against much of the uncertainty generated in incorrect structural parameters, provided the 483 satellite-derived LAI value is reasonable. Underestimations of the actual ground LAI value by the satellite 484 product may therefore cause substantial uncertainty in modelled Chl_{Leaf} results.

- 485
- 486

2.5 Data smoothing and gap-filling

487 The Locally Adjusted by Cubic-Spline Capping (LACC) method (Chen, Deng, and Chen 2006) was 488 developed to produce continuous seasonal trajectories of satellite-derived surface parameters 489 contaminated by clouds and other atmospheric effects. The LACC method previously has been used 490 successfully to smooth time series of clumping index and NDVI (He et al. 2016). Contaminated points can 491 be identified in the time series and replaced with expected values through temporal interpolation 492 between adjacent valid points. For identifying atmospherically contaminated points in a time series, a 493 cubic-spline curve fitting technique with a curvature constraint was developed. Without this curvature 494 constraint, the fitted polynomial curve would also fit the contaminated points, defeating the study 495 purpose. In the LACC algorithm, a maximum global curvature of 0.5 is prescribed in the first step to 496 identify contaminated points. In the second step, the curvature is adjusted locally (i.e. at different dates 497 of the year) according to the local shape of the curve fitted with the global curvature constraint. This 498 adjustment is to enhance the large curvatures generally found at the beginning and the end of a growing 499 season, or double cropping seasons, and to reduce the curvature during a non-growing season and/or 500 around the peak of growing seasons. In this way, smooth trajectories that also follow the rapid growing 501 season variation patterns can be produced. In this study, the LACC algorithm is used to identify

502 contaminated data points and to gap-fill missing data points. However, in tropical areas there are often 503 fewer than the required six data points needed to run the LACC algorithm. In this case, in order to derive 504 temporally continuous data across the year, we removed the highest and lowest data point and linearly 505 interpolated across the year. In northern latitude evergreen needleleaf forests where snow cover 506 prevented acquisition of surface reflectance data from the vegetation canopy, we took the mean 507 seasonal value and extended it across the year. Whilst such temporal gap-filling is likely to introduce 508 some uncertainty in the modelled chlorophyll results, the temporal variations in chlorophyll content for 509 these PFTs is small relative to deciduous forests or croplands. The annual global distribution of leaf 510 chlorophyll content maps that are presented in this paper are for the last full year that MERIS was 511 operational (2011). Missing data due to cloud or MERIS image acquisition limitations (Tum et al. 2016) 512 are gap-filled with corresponding 2010 data from the same DOY within the LACC smoothing algorithm 513 for spatial completeness and visual assessment. The original 2003-2011 global maps are also available 514 for use within the academic community.

515

516 3.0 Results and Discussion

517 The annual global distribution of leaf chlorophyll content is presented in Figure 4a, with the geographic 518 trends varying both within and between biomes. Figure 4b depicts the annual range in chlorophyll 519 values for the major PFTs, as an integration of the temporal and spatial variability in chlorophyll content 520 over a year and across the globe.





522

Figure 4: a) The spatial distribution of median annual global distribution of leaf chlorophyll content (at 9
 km resolution), and b) the median and interquartile range of leaf chlorophyll content, along with
 extreme values given for 9% and 91% (whiskers) and 2% and 98% (circles) of the data range.

527 On an annual basis, the highest median Chl_{Leaf} values are present in evergreen broadleaf forests (54.4 μ g 528 cm^{-2} ; Figure 4b), which is in part due to the high chlorophyll content of the vegetation present, and also 529 due to the lack of seasonal leaf loss. The lower median annual chlorophyll values (Figure 4b) are consequently found in deciduous biomes (broadleaf and needleleaf forests, 28.8 µg cm⁻² and 11.4 µg 530 531 cm², respectively). Ground measurements have previously demonstrated that nitrogen-poor needleleaf 532 species typically exhibit lower chlorophyll contents by unit area than broadleaf species (Middleton et al. 533 1997), which is also evident in the median annual results shown in Figure 4 (ENF, 18.9 μ g cm⁻²). 534 Environmental constraints may also impact Chl_{Leaf} values, through temperature extremes or water 535 availability in evergreen PFTs. Temperate grasslands, for example, are often located in regions that 536 experience extreme annual temperature variability (McGinn 2010), which affects chlorophyll 537 biosynthesis (Ashraf and Harris 2013). From the global map of median annual Chl_{Leaf}, the footprints of

538 some agricultural regions are also clearly detectable, for example the 'Corn Belt' in Midwestern USA,

539 throughout India, the Loess belt in central Europe and in northeastern China, due to their higher annual

540 peak values, which may be related to fertilization application.

541

542 **3.1** Validation of modelled leaf chlorophyll estimates for individual PFTs

The leaf chlorophyll algorithm was validated using 248 samples, across a number of different sampling dates and years. The modelled chlorophyll results are plotted against ground-measured values for all available PFTs (Figure 5).

546



Figure 5: Validation of modelled leaf chlorophyll with measured ground data for individual plant functional types. Data are distinguished between temporally matched sampling and satellite overpass dates, and for ground data collected outside the MERIS lifespan (2002-2012), the closest satellite date was selected.

The strongest performances are seen for DBF ($R^2 = 0.67$; RMSE = 9.25 µg cm⁻², relative RMSE = 25.4%) 554 555 followed by ENF ($R^2 = 0.47$; RMSE = 10.63 µg cm⁻², relative RMSE = 32.62%) and CRP ($R^2 = 0.41$; RMSE = 13.18 μ g cm⁻², relative RMSE = 30.8%). Nonetheless the other three PFTS also showed similar levels of 556 557 uncertainty, where relative RMSE values were 26.5%, 15.7% and 20.8% for EBF, GRS and SHR, 558 respectively. The statistics reported in Figure 5 are for combined temporally matched sampling and satellite overpass dates, and for ground data collected outside the MERIS lifespan (2002-2012), where 559 560 the closest satellite date was selected (See Section 2.1). For comparison, the regression results using only the matched dates inside the MERIS operational window are: DBF: $R^2 = 0.63$, p<0.001; ENF: $R^2 =$ 561 562 0.63, p<0.001; EBF: insufficient data; and CRP: $R^2 = 0.43$, p<0.001. The shrub data used in Figure 5 only contained matched dates. The regression results for all dates combined and only matched dates are 563 564 comparable, with the largest difference arising for EBF, which has fewer data and a smaller dynamic 565 range.

566

567 **3.2** Overall validation and comparison with vegetation indices

The transferability of an algorithm across spatial and temporal scales is essential for modelling Chl_{Leaf}, or 568 569 any ecological variable, at the global scale. In comparison to empirical approaches (Gitelson, Gritz, and 570 Merzlyak 2003, Roberts, Roth, and Perroy 2016, Peng et al. 2017), the nature of physically-based 571 retrieval methods can account for relationships between canopy reflectance and Chl_{Leaf} across different species and measurement acquisition conditions. Regressions between measured and modelled ChlLeaf 572 573 for all PFTs combined are shown in Figure 6, alongside results for two popular vegetation indices with the same measured Chl_{Leaf} data (the chlorophyll-sensitive MERIS Terrestrial Chlorophyll Index (MTCI = 574 575 754 nm-709 nm/709 nm-681 nm) and the Normalised Difference Vegetation Index (NDVI = 865 nm-664 576 nm/865 nm+664nm)).



Vegetation Index (NDVI) values.

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583

Figure 6a demonstrates the suitability of the Chl_{Leaf} algorithm for application across multiple PFTs, with a 584 strong, linear relationship for all PFTs combined (R² = 0.47; p<0.001). By contrast, the MTCI regression is 585 586 weaker (R² = 0.27; p<0.001), with some separation in MTCI values according to PFT. Cropland and DBF, 587 for example, exhibit higher values than grassland and shrubland, for the same ChlLeaf. This stratification is 588 likely to due to the strong influence of LAI on MTCI values. Nonetheless, MTCI does present a relative 589 improvement over NDVI results ($R^2 = 0.02$; n/s), due to the inclusion of chlorophyll-sensitive red-edge 590 bands within the VI. The biomass-sensitive NDVI values also exhibit a separation according to PFT, due 591 to differences in canopy structure and background, with cropland and grasslands also presenting 592 particular variability in values within the same PFT. This result points to the importance of accounting for 593 variations in canopy structure when deriving leaf-level chlorophyll results, and has implications for 594 applying VIs over large spatial extents to infer information on physiological processes or plant 595 productivity.

596

597 **3.4 Leaf chlorophyll phenology and temporal trends**

598 The annual variability in leaf chlorophyll content was evaluated through the standard deviation of mean 599 annual Chl_{Leaf}, (Figure 7a). The seasonal phenology of mean Chl_{Leaf} and mean LAI across the northern 600 hemisphere are shown for different biomes in Figure 7b.



602

Figure 7: Global 2011 maps of a) seasonal variability in leaf chlorophyll (one standard deviation of one
 year time series), and b) mean seasonal phenologies of modelled chlorophyll content and LAI across the
 northern hemisphere for six different plant functional types during 2011. Shaded area indicates one
 standard deviation.

607

Global regions that contain large areas of deciduous forests and croplands, which are dominated by a strong seasonal phenology, present the largest temporal variation and dynamic range in Chl_{Leaf} values (Figure 7a). In Figure 7b, DBF displays an expected phenology associated with budburst and chlorophyll biosynthesis in spring (from circa DOY 120) and chlorophyll breakdown during leaf senescence. 612 Temporal variations in cropland Chl_{Leaf} show a high variability globally, which is associated with different 613 species of crops planted, planting regimes (i.e. single or double), fertiliser application and the level of 614 irrigation. ENF temporal Chl_{Leaf} trajectories are relatively consistent across the year, although increasing Chl_{Leaf} within new needles in spring is detectible, along with some chlorophyll breakdown within winter 615 months. Missing local data (predominately in the Amazon) within Figure 7a is due to too few original 616 617 data points to generate standard deviation values. Importantly, Figure 7b demonstrates the temporal 618 divergence of LAI and Chl_{Leaf} across the growing season, and a clear decoupling of vegetation structure 619 and physiological function (Croft, Chen, and Zhang 2014b, Walther et al. 2016).

620

621 **3.3 Spatial and biome-dependent trends**

Annual maximum chlorophyll maps offer an opportunity to examine spatial differences in Chl_{Leaf} without the integration of a temporal bias from seasonal change. The geographic variability in the abundance of leaf chlorophyll depends in part on local environmental drivers, and partly on plant resource allocation.

626



Figure 8: a) Maximum annual leaf chlorophyll content (at 9 km resolution) and b) mean values of the
maximum along latitudinal bands.

632

633 Higher annual maximum chlorophyll values (> 55 μ g cm⁻²) are associated with tropical forests and croplands (Figure 8a). Mid-range values (25-55 μ g cm⁻²) are typical of grasslands, temperate broadleaf 634 forests and arctic tundra vegetation, and lower values (<25 μ g cm⁻²) are typical of boreal forests. 635 636 Considerable intra-biome chlorophyll variability also exists, for example within croplands higher annual 637 values are present in India, Eastern China, Western Europe and the great Lakes in USA/Canada, in 638 contrast to lower values found in Central Europe and the North American mid-west. Reasons for 639 chlorophyll variability over space are in part due to the abundance of elements needed for chlorophyll 640 synthesis (nitrogen (N), phosphorous (P), Magnesium (Mg)) (Li et al. 2018), which may vary due to the 641 nutrient availability status of the soil, the addition of fertilisers in managed landscapes and as a result of 642 atmospheric N deposition. At the biome-scale, the temperature-dependency of chlorophyll synthesis 643 reactions, where the optimum temperature of [3,8-DV]-Pchlide a 8-vinyl reductase (DVR) enzyme activity in chlorophyll synthesis is 30°C (Nagata, Tanaka, and Tanaka 2007), may exert a biotic control on 644

the accumulation of chlorophyll in temperature-limiting ecosystems, such as the arctic. Additionally, plant water availability and drought stress affect the synthesis of chlorophyll, and may also prompt its accelerated breakdown (Ashraf and Harris 2013). As plants allocate resources to optimise physiological processes, it may be expected that solar irradiance conditions, and the leaf economics spectrum (Wright et al. 2004) affect the partitioning of nitrogen, both between and within structural and photosynthetic leaf fractions (Croft et al. 2017, Niinemets and Tenhunen 1997).

651

4.0 Sources of uncertainty in modelled leaf chlorophyll content

653 The validation results (Figure 5 and 6a) indicate a good algorithm performance and levels of uncertainty 654 comparable to more local scale studies, with reported RMSE values in the literature ranging from around 8 to 15 μ g cm⁻² (Houborg et al. 2015). The uncertainties values from all of the PFTs used in this study 655 were within this range; from GRS: 5.71 μ g cm⁻² through to EBF: 14.94 μ g cm⁻², and the overall 656 657 uncertainty of all PFTs considered together was 10.81 µg cm⁻². The overall uncertainty of the global 658 Chl_{Leaf} product is therefore considered within reasonable limits of current state of the art approaches to 659 chlorophyll modelling. The sources of uncertainty and future ways to minimise this uncertainty are 660 discussed further below.

661

We highlight the main sources of uncertainty that can affect the accuracy of the modelled leaf chlorophyll values within the global maps as: 1) the data quality of the MERIS surface reflectance, 2) the accuracy of input LAI data, 3) the use of fixed structural parameters in canopy models, 4) the spectral contributions of understory or background reflectance to satellite-derived canopy reflectance, and 5) the validation sites used within the study.

667

1) Uncertainty in the MERIS surface reflectance product can arise from sources such as sensor calibration issues, cloud contamination, atmospheric correction errors (Garrigues et al. 2008). Due to the nature of the inversion process and the 'ill-posed problem' (Combal, Baret, and Weiss 2002), even small changes in satellite-derived reflectance can lead to large variations in modelled chlorophyll values (Garrigues et al. 2008).

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674 2) The chlorophyll retrieval algorithm is strongly reliant on the LAI as an input parameter, where small 675 errors in LAI values, particularly at low LAI values can lead to large errors in modelled chlorophyll. This 676 introduces a greater uncertainty into the chlorophyll estimates in areas with little vegetation cover, for 677 example in shrubland and deciduous plant forms at the start and end of the growing season. Errors in 678 LAI at high LAI values, for example due to problems concerning reflectance saturation are less of a 679 concern because the algorithm is less responsive to LAI > 4, or when the canopy behaves as a 'big leaf' 680 and background and branch contributions are minimal. Clumping is considered in the GEOV1 LAI 681 product that we selected (Baret et al., 2013), at both the landscape scale and the canopy scale through 682 the integration of CYCLOPES v3.1 and MODIS c5 biophysical products. Errors in LAI values that arise from 683 a lack of proper consideration of clumping effects mainly affect needleleaf canopies, which can be 684 largely underestimated (Chen, Menges, and Leblanc 2005). The most significant problem, however, for 685 the modelling of Chl_{Leaf} is a lack of accurate LAI during winter months in the northern hemisphere. The 686 presence of snow on the understory and on the actual needleleaf shoots prevents the retrieval of LAI 687 during large periods of time in the winter months. Any satellite-derived LAI data that is retrieved during 688 snow-dominated winter months is highly uncertain, leading to potential problems with the smoothing 689 algorithm. Further uncertainty arises from the LAI data product's coarser spatial resolution (1000 m), 690 compared to the input MERIS surface reflectance product (300 m). The greatest uncertainty arising from 691 this source will be in patchy or spatially variable landscapes, such as croplands or where the LAI values are low (i.e. LAI = <3). This uncertainty will have less of a bearing on more spatially homogenous
landscapes (i.e. broadleaf forests) and in vegetation with higher LAI values.

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695 3) The chlorophyll retrieval algorithm considers the major PFTs separately, through the application of 696 individual LUTs based on measured and reported ground data. However, the structural values used to 697 create the LUTs represent a generalised approximation of the structural parameters, and it is recognised 698 that considerable variation can exist within a given biome. An improved spatial representation of the 699 intra-PFT variability of structural parameters, such as canopy height, stem density and leaf angle 700 distribution, would be beneficial, particularly in needleleaf forests, for example, where spatial variability 701 is driven by the dominant species composition. However, a lack of available spatially-continuous data at 702 fine spatial resolutions currently prevents the explicit parameterisation of these structural variables in 703 the canopy models. Whilst these structural parameterisations are important, their variation over the 704 variable range that may be found within a PFT is likely to be small relative to variations between PFTs, 705 and relative to LAI. The sensitivity analysis presented in Section 2.4 indicates that LAI is the dominant 706 driver of modelled Chl_{Leaf} uncertainty, with other parameters such as crown height, ΩE , γE , Hs parameter 707 the background soil factor having negligible influence. This finding also confirms the work of Zhang et al. 708 (2008), who examined the sensitivity of the 4-Scale model input parameters to the model output, 709 finding that LAI is the dominant driver of modelled canopy reflectance.

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4) The temporal and spatial variability of background contributions are a source of uncertainty in the modelled Chl_{Leaf} values, to a greater or lesser degree depending on the overall canopy coverage. Areas with lower LAI values are more susceptible to errors concerning the erroneous parameterisation of background material in the model. Although substantial progress has been made in retrieving understory reflectance from multi-angular reflectance images (Canisius and Chen 2007, Pisek and Chen

2009), the often coarse spatial resolution and lack of global coverage for all vegetation types of these
products (Jiao et al. 2014, Liu et al. 2017) precludes their use.

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719 5) An additional source of uncertainty could come from the validation ground data itself. The relatively 720 small number of measured chlorophyll values sampled in situ makes widespread validation challenging, 721 particularly for EBF, SHR and DNF PFTs. The validation data used in this study is through the generous 722 and collaborative efforts of independent researchers, and without an established network collecting 723 regular Chl_{Leaf} data or automated measurement systems, direct product validation is time and resource 724 intensive (Garrigues et al. 2008). Consequently, no existing validation data sets are completely 725 representative of all of the global and seasonal variability of vegetation [Baret et al., 2006]. This study 726 used 28 distinct sites to validate the Chl_{leaf} product, which is within the same range of other global 727 biophysical product validations. In the calibration and product development of MODIS LAI products, 728 Yang et al. (2006) selected 25 validation sites, and He et al (2012) used 38 validation sites for their 500 m 729 global clumping index map. Initial Soil Moisture Active Passive (SMAP) soil moisture products were 730 validated using 34 validation sites (Colliander et al. 2017). To an extent, the reported uncertainties on 731 the modelled leaf chlorophyll content values will also be a function of the coverage of geographic and 732 vegetation species contained within the validation sites, and the inherent variability that exists within 733 plant functional types. Future work can focus on evaluating the accuracy of the chlorophyll product in 734 species and management regimes that are underrepresented in the model development and validation. 735 It is likely that sites where the structural properties deviate largest form the values used in the model 736 parameterisation (Table 2) may suffer the largest degree of uncertainty.

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740 **5.0 Conclusion**

741 This research represents the first global view of terrestrial Chl_{Leaf} distribution. Weekly maps of Chl_{Leaf} are 742 produced at the global scale following a two-step physically-based modelling approach. The accuracy of 743 the Chl_{Leaf} product is ultimately dependent on the representation of radiative transfer processes within 744 the canopy and leaf optical models, the structural parameterisation of the radiative transfer models and 745 the accuracy of land cover and leaf area index data. Modelled results show good relationships with measured ground data, in particular for deciduous broadleaf forests ($R^2 = 0.67$; RMSE = 9.25 µg cm⁻²; 746 p<0.001), croplands (R² = 0.41; RMSE = 13.18 μ g cm⁻²; p<0.001) and evergreen needleleaf forests (R² = 747 0.47; RMSE = 10.63 μ g cm⁻²; p<0.001). On an annual basis, evergreen broadleaf forests presented the 748 749 highest median leaf chlorophyll values (54.4 µg cm⁻²). The global values show large temporal and spatial 750 variability expected in Chl_{Leaf}. It is expected that ESA Sentinel-2 series will be used to continue the Chl_{Leaf} 751 time series, due to the presence of red-edge sampling bands and widespread spatial coverage and fine 752 temporal resolution. It is anticipated that this global leaf chlorophyll product will make a significant step 753 towards improving global and regional ecosystem models associated with carbon cycle modelling, 754 through the explicit consideration of foliage biochemistry, and reduce the uncertainty associated with 755 leaf physiology.

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764 7.0 References

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