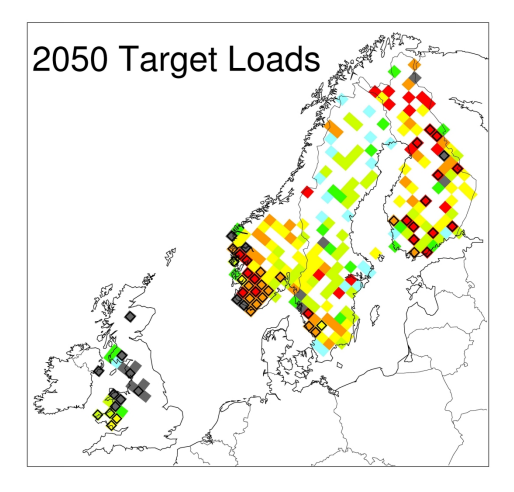
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Maximilian Posch, Julian Aherne, Filip Moldan, Chris D. Evans, Martin Forsius, Thorjørn Larssen, Rachel Helliwell, and B. Jack Cosby. Dynamic Modeling and Target Loads of Sulfur and Nitrogen for Surface Waters in Finland, Norway, Sweden, and the United Kingdom. Environmental Science & Technology. 2019, 53(9), 5062-5070.



TOC Figure 193x186mm (300 x 300 DPI)

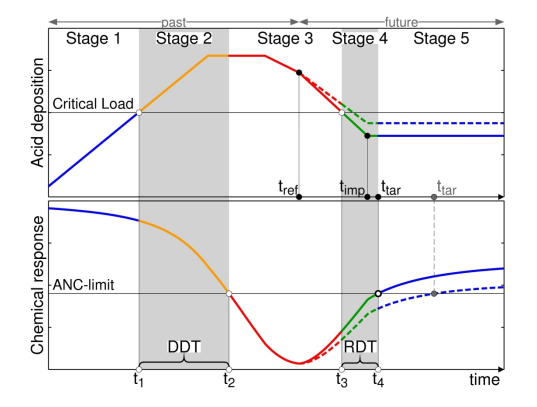


Figure 1: Generalised past and future development stages (indicated by different colours) of acidic deposition and lake chemical criterion response (here: ANC) in comparison to the critical chemical value (ANC-limit) and the critical load derived from it (i.e., the determination of critical load is based on a critical limit for a specified chemical criterion, which protects the biological indicator from deleterious effects). The delay between the (non-)exceedance of the critical load and the (non-)violation of the critical chemical value is indicated in grey shades, highlighting the Damage Delay Time (DDT) and the Recovery Delay Time (RDT) of the system. Also shown are the points in time (tref, timp) relevant for defining a target load (< critical load) to reach non-violation of the chemical value at a pre-specified time ttar. The dashed lines show the temporal development for a later target year (labelled in grey).

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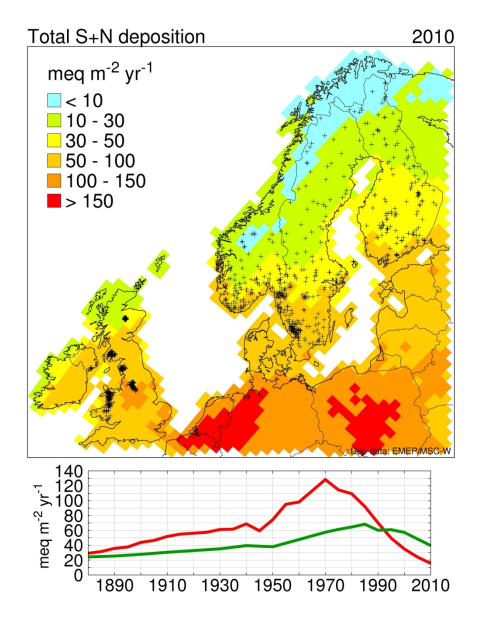


Figure 2: Location of the study sites used for dynamic modelling (n = 848; black crosses) overlaid upon modelled total sulphur (S) plus nitrogen (N) deposition in 2010 (meq m-2 yr-1) on the EMEP 50 km × 50 km grid (for further details on the EMEP model see Simpson et al.57). The temporal development (1880–2010) of the total S (red) and N (green) deposition averaged over the 848 study sites is also shown.

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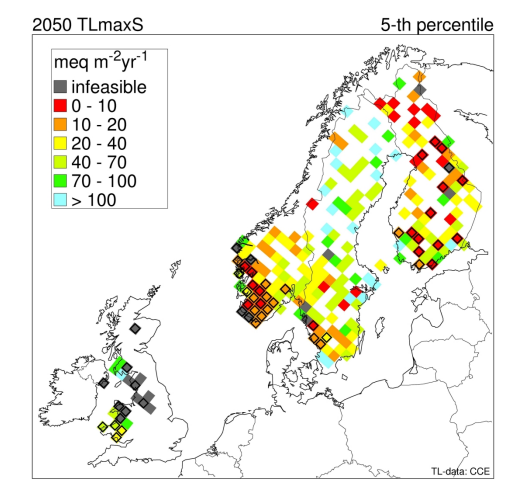


Figure 3: Map of the 5-th percentile of the 2050 maximum target load of sulphur (TLmaxS) on the EMEP 50 km × 50 km grid for 848 catchments in Finland, Norway, Sweden and the United Kingdom. Note: TLmaxS was set to the maximum critical load of S (CLmaxS) where TLmaxS > CLmaxS (i.e., the critical load is sufficient for non-violation of the ANC-limit by 2050). Grey-filled cells (label `infeasible') denote grids containing at least one lake where the simulated ANC does not meet the specified limit by 2050, even under zero deposition after 2020 (FI = 3, NO = 4, SE = 4, UK = 35; Table 1). Black diamonds frame grids with at least one `true' TL, i.e., where a TL exists and is lower than the CL (FI = 21, NO = 52, SE = 10, UK = 44; Table 1).

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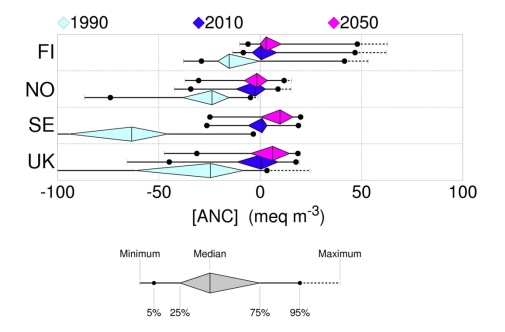


Figure 4: Percentile statistics ('diamond plots') of simulated annual average lake acid neutralising capacity (ANC) in 1990, 2010 and 2050 in Finland (FI), Norway (NO), Sweden (SE) and the United Kingdom (UK). Data are only shown for lakes for which target loads were determined (FI = 24, NO = 56, SE = 14, UK = 79; see columns 'INF' plus 'TL' in Table 1).

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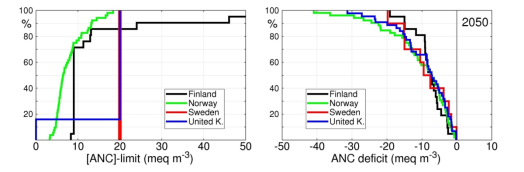


Figure 5: Left: Cumulative distribution functions of the ANC-limits for the lakes with target loads in Finland (21), Norway (52), Sweden (10) and the United Kingdom (44). Right: Inverse cumulative distribution functions of the 'ANC deficit' in 2050 for the same lakes in those countries. The ANC deficit shows the difference between simulated ANC in 2050 under the 2010 Gothenburg Protocol (see Figure 2) and the specified critical ANC-limit (left Figure).

152x50mm (300 x 300 DPI)

1	Dynamic Modelling and Target Loads of Sulphur and Nitrogen for Surface Waters in
2	Finland, Norway, Sweden and the United Kingdom
3	
4	
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18	
19	
20	Abstract
21	The target load concept is an extension of the critical load concept of air pollution inputs to
22	ecosystems. The advantage of target loads over critical loads is that one can define the
23	deposition and the point in time (target year) when the critical (chemical) limit is no longer
24	violated. This information on the timing of recovery requires dynamic modelling. Using a
25	well-documented dynamic model, target loads for acidic deposition were determined for 848
26	surface waters across Finland, Norway, Sweden and the United Kingdom for the target year
27	2050. In the majority of sites ($n = 675$), the critical ANC-limit was predicted to be achieved
28	by 2050; however, for 127 sites target loads were determined. In addition, 46 sites were
29	infeasible, i.e., even a deposition reduction to zero would not achieve the limit by 2050. The
30	average maximum target load for sulphur was 38% lower than the respective critical load
31	across the study lakes ($n = 127$). Target loads on a large regional scale can inform effects-
32	based emission reduction policies; the current assessment suggests that reductions beyond the
33	Gothenburg Protocol are required to ensure surface water recovery from acidification by
34	2050.
35	

Keywords. Acidification; critical load; recovery; Acid Neutralising Capacity (ANC); Model
 of Acidification of Groundwaters in Catchments (MAGIC)

38 39

40 **1. Introduction**

41 During the 1970s it was recognized that surface waters in large parts of Europe and North 42 America were being influenced by air pollution, i.e., acidic deposition, owing to anthropogenic emissions of sulphur (S) and nitrogen (N) oxides.^{1,2} Shortly thereafter, 43 44 empirical³ and steady-state⁴⁻⁶ models were developed and applied to predict the impacts of 45 acidic deposition on surface waters. In concert, it was recognised that time-dependent 46 processes could buffer (delay) ecosystem (soil and surface water) response to acidic 47 deposition. The incorporation of these processes required time-dependent or 'dynamic' 48 modelling frameworks. The earliest dynamic simulation models incorporated established 49 relationships from soil and water chemistry to predict the most likely effects of acidic 50 deposition on surface waters.⁷ Ultimately, these dynamic models provided a quantitative 51 framework to assess whether (and how quickly) a decrease in acidic stress would result in a recovery of ecosystems.⁸ Since the 1980s, several dynamic (hydro-chemical) models have 52 53 been developed and extensively applied at site-specific and regional scales to predict changes 54 in soil and surface water chemistry due to acidic deposition.⁹⁻¹⁵ Moreover, dynamic models 55 can provide a quantitative estimate of the time lag between a reduction in deposition and the 56 attainment of 'acceptable' ecosystem status (based on a threshold, or 'critical value', for a 57 specified chemical criterion, e.g., surface water pH=6.0). This time lag has been denoted as the damage time lag¹⁶ or recovery delay time.¹⁷⁻¹⁹ 58

59

60 The assessment of impacts of acidic deposition on terrestrial and aquatic ecosystems has 61 supported policies to reduce anthropogenic S and N emissions. In Europe, the critical loads 62 approach is widely accepted as the basis for negotiating effects-based control strategies for air pollution. A critical load is defined as 'a quantitative estimate of an exposure to one or more 63 64 pollutants below which significant harmful effects on specified sensitive elements of the environment do not occur according to present knowledge'.²⁰ The approach is based on 65 setting a critical limit for a chemical criterion (e.g., Acid Neutralising Capacity [ANC]) to 66 67 protect a specified biological indicator for a chosen receptor ecosystem (e.g., fish species for 68 surface waters, or tree roots in forest soils), and via inverse modelling a deposition (the 69 critical load) is derived to ensure the limit is not violated and thus 'harmful effects' avoided.²¹⁻²³ Critical loads have been used in the negotiations of several protocols to the 70 71 United Nations Economic Commission for Europe's (UNECE²⁴) Convention on Long-range

Transboundary Air Pollution (LRTAP) and the European Union's National Emission Ceilings
 (NEC) Directive²⁵ and its revision.

74

75 However, by definition, critical loads are steady-state quantities and as such they do not 76 provide information on the time involved for a system (e.g., surface water) to reach a certain 77 chemical (or biological) state. Dynamic models are needed to calculate recovery times under 78 prescribed emission reductions. As such, dynamic modelling has also become an important 79 part of the effects-oriented work under the LRTAP Convention.²⁶ If a desired chemical state of a surface water is defined for a given year, dynamic models can be used in an inverse mode 80 81 to compute the deposition path leading to that desired state (if feasible). Such a deposition is called a target load, and the year in which the desired state is to be reached is called the target 82 83 year.^{18,19} There have been few published studies describing the use of target loads in Europe^{16,17,27} and fewer for surface waters.²⁸⁻³⁰ It is important to note that in North America 84 85 the term 'target load' has also been used to refer to a 'target' deposition, determined by 86 political (or management) agreement, that can be higher or lower than the critical load³¹⁻³⁵, 87 often based on arbitrary interpretations of the impacts data rather than the avoidance of 88 specific deleterious ecological effects.³² 89

90 The objective of this study was to establish target loads for European regions dominated by

91 acid-sensitive surface waters, which ensure acidification recovery by the year 2050 (target

92 year). The target loads go beyond deposition reductions under the Gothenburg Protocol

93 (implemented in 2010), to ensure chemical recovery in surface water ANC (chemical

94 criterion). The target loads were determined using the Model of Acidification of

95 Groundwaters in Catchments (MAGIC) for lakes in Finland, Norway, Sweden, and the United

Kingdom. Further, the conceptual basis for the determination of a target load from a dynamicmodel is also provided.

98

99 2. Dynamic Modelling and Target Loads

100 With critical loads, i.e., in the steady-state situation, only two cases can be distinguished when 101 evaluated against deposition: (1) deposition is below the critical load, or (2) deposition is 102 greater than the critical load, i.e., there is an exceedance of the critical load. In the first case 103 there is no (apparent) problem, i.e., no reduction in deposition is deemed necessary. In the 104 second case there is, by definition, an increased risk of damage to the ecosystem. Thus, a 105 critical load serves as a warning as long as there is exceedance, since it indicates that 106 deposition should be reduced. However, it is often (implicitly) assumed that reducing 107 deposition to (or below) the critical load immediately removes the risk of 'harmful effects', 108 i.e., the critical chemical criterion (e.g., the ANC-limit) that links the critical load to the

109 (biological) effect, immediately attains a non-critical ('safe') value (and that there is

- 110 immediate biological recovery as well). However, the reaction of an ecosystem (or
- 111 catchment) to changes in deposition is delayed by (finite) buffers, e.g., the cation exchange
- 112 capacity of catchment soils. These buffer mechanisms can delay the attainment of the critical
- 113 chemical value, and it might take decades or even centuries, before steady state is reached.
- 114 These finite buffers are not included in the critical load formulation, since they do not
- 115 influence the steady state, but only the time to reach it.
- 116

117 Therefore, dynamic models are needed to estimate the time involved in attaining a certain

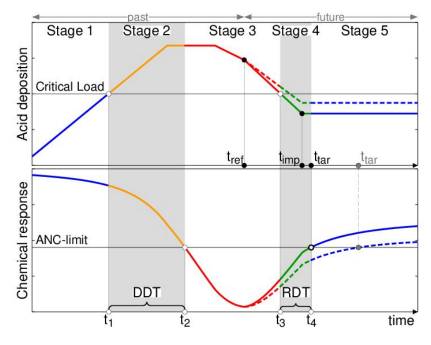
- 118 chemical state in response to different deposition scenarios. Five stages can be distinguished
- in the possible development of a (lake) chemical variable in response to a 'typical' temporal
- 120 deposition pattern (see Figure 1):
- 121

Stage 1: Deposition is below the critical load (CL) and the chemical criterion (ANC) does not
violate its critical limit. As long as deposition stays below the CL, this is the 'ideal' situation
(blue lines in Figure 1).

- 125 Stage 2: Deposition is above the CL, but the critical chemical criterion is not (yet) violated
- 126 because there is a time delay of ecosystem response before adverse effects occur. Therefore,
- 127 no damage is likely to occur at this stage, despite exceedance of the CL. The time between the
- 128 first exceedance of the CL and the first violation of the chemical criterion is termed the
- 129 *Damage Delay Time* (DDT= t_2-t_1).
- 130 *Stage 3*: The deposition is above the CL and the critical chemical criterion is violated.
- 131 Measures (emission reductions) have to be taken to avoid a (further) deterioration of the
- 132 ecosystem (biological indicator linked to the chemical criterion).
- 133 Stage 4: Deposition is below the CL, but the chemical criterion is still violated and thus (full)
- 134 recovery has not yet occurred. The time between the first non-exceedance of the CL and the
- 135 subsequent non-violation of the criterion is termed the *Recovery Delay Time* (RDT= t_4-t_3).
- 136 Note: RDT is not necessarily the same (or even similar) to DDT due to hysteresis effects in
- 137 certain (soil) processes (e.g., cation exchange); the schematic is purely for illustration and
- 138 does not reflect the relative temporal processes.
- 139 *Stage 5*: Deposition is below the CL and the critical chemical criterion is no longer violated.
- 140 This stage is analogous to Stage 1, and the ecosystem is considered to have recovered. In
- 141 practice it might happen that the critical limit cannot be achieved within a reasonable (policy-
- 142 relevant) timeframe, even for zero N and S deposition, e.g., due to the depletion of
- 143 exchangeable base cations.

144

- 145 In addition to the delay in chemical recovery, there is likely to be a further delay before the
- 146 'original' biological state is reached, i.e., even if the chemical criterion is met, it will take
- 147 time before biological recovery is achieved (if at all).
- 148



149

150 Figure 1: Generalised past and future development stages (indicated by different colours) of acidic 151 deposition and lake chemical criterion response (here: ANC) in comparison to the critical chemical 152 value (ANC-limit) and the critical load derived from it (i.e., the determination of critical load is based 153 on a critical limit for a specified chemical criterion, which protects the biological indicator from 154 deleterious effects). The delay between the (non-)exceedance of the critical load and the (non-155)violation of the critical chemical value is indicated in grey shades, highlighting the Damage Delay 156 Time (DDT) and the Recovery Delay Time (RDT) of the system. Also shown are the points in time 157 (t_{ref}, t_{imp}) relevant for defining a target load (< critical load) to reach non-violation of the chemical value 158 at a pre-specified time t_{tar}. The dashed lines show the temporal development for a later target year 159 (labelled in grey).

160

161 The most straightforward use of a dynamic model is scenario analysis, i.e., first a future 162 deposition scenario is assumed, and then the (chemical) consequences for the ecosystem (e.g., 163 lake) are evaluated. A target load, on the other hand, is the deposition path that ensures that a 164 prescribed value of the chemical criterion is achieved in a given year. Here we define a target 165 load as a deposition path characterised by three numbers (years): (i) the reference year, (ii) the 166 implementation year, and (iii) the target year (see Figure 1). The *reference year*, t_{ref} , is the year (time) up to which the (historic) deposition path is given and cannot be changed. The 167 168 implementation year, timp, is the year by which all reduction measures, needed to reach the 169 final deposition (the target load), are implemented. Between the reference year and the

170 implementation year depositions are assumed (assured) to change linearly (see Figure 1). 171 Finally, the *target year* is the year in which the critical chemical criterion (e.g., the ANC-172 limit) is met (for the first time). The above three years define a unique deposition path, the 173 final value of which is referred to as a target load. The earlier the target year, the lower the 174 target load (at sites where the chemical criterion is violated – for other sites a target load is 175 not relevant), since higher deposition reductions are needed to achieve the desired status 176 within a shorter timeframe (see Figure 1). In extreme cases, a target load might not exist at 177 all, i.e., even reduction to zero deposition in the implementation year will not result in the 178 desired ecosystem status within the prescribed time; in this instance the target load is termed 179 'infeasible'. For more information on target loads and related topics see Posch et al.²², Jenkins et al.¹⁹ or Bonten et al.²⁷ 180

181

182 **3. Materials and Methods**

183 The current study focused on surface waters (lakes and streams) with long-term observations 184 of chemistry in acid-sensitive regions of Europe³⁶, i.e., Finland, Norway, Sweden, and the 185 United Kingdom. In general, these sites are considered to be sentinel indicators of 186 acidification impacts, and their recovery is assumed to reflect wider ecosystem acidification 187 recovery across the entire study region; as such, they are well suited for the determination of 188 regionally representative target loads. All surface waters were part of routine acidification 189 monitoring networks since the 1980s and 1990s, typically located in regions with acid-190 sensitive geology. For example, the Finnish acidification monitoring network, maintained by 191 the Finnish Environment Institute, consisted of 163 lakes located throughout Finland, 192 subjectively chosen by expert judgement from a national acidification survey during 1987 for 193 use in acidification assessments.³⁷ Similarly, the Norwegian study lakes (n = 131) were a 194 subset of the national monitoring programme, confined to lakes south of 62.5° latitude, with 195 observations suitable for dynamic modelling. The study sites are predominantly small acid-196 sensitive headwater lakes and streams, with low base cation concentrations, low alkalinity and low (charge balance) ANC.³⁸ All surface waters have been widely used in acidification 197 198 assessments evaluating long-term trends in surface water chemistry³⁹⁻⁴¹, and the prediction of 199 future chemistry using dynamic (hydro-chemical) models, specifically MAGIC.⁴²⁻⁴⁶ The study 200 sites have played a central role in European-scale projects, such as 'Recover:2010'47 and 201 'Eurolimpacs'⁴⁸, focused on model simulations of surface water response to European 202 emissions reduction policies. The process limitations and predictive uncertainty of MAGIC in 203 isolation, and compared with other models, e.g., PnET-BGC (photosynthesis and 204 evapotranspiration-biogeochemistry), SAFE (soil acidification in forest ecosystems), and VSD (very simple dynamic), have been widely published.^{12,15,49,50} Similarly, the influence of 205 climate change on model predictions for MAGIC have been widely assessed^{43,46,51,52}. As such, 206

herein we focus on the determination of target loads using MAGIC, which (hitherto for) have
not been reported for the study sites and refer the reader to previous publications for detailed
information regarding model calibration and process uncertainty for MAGIC. Nonetheless,
we provide a brief overview of MAGIC, its application, calibration and simulation for the
study sites, specifically with respect to target loads.

212

213 MAGIC is a lumped-parameter model of intermediate complexity, developed to predict the 214 long-term effects of acidic deposition on soil and surface water chemistry.^{53,54} The model 215 predicts monthly and annual average concentrations of the major ions for soil solution and 216 surface water chemistry. MAGIC represents the catchment with aggregated, uniform soil 217 compartments (up to three), and a surface water compartment that can be either a lake or a 218 stream. Time series inputs to the model include: deposition of ions from the atmosphere (wet 219 plus dry deposition); discharge volume and flow routing within the catchment; biological 220 production and removal of ions; internal sources and sinks of ions from weathering or 221 precipitation reactions; and climate data. Constant parameters in the model include physical 222 and chemical characteristics of the soils and surface waters, and thermodynamic constants. 223 Soil base cation weathering rate and initial base saturation are calibrated using observed 224 values of surface water and soil chemistry for a specified period. In this instance, calibration 225 refers to an automated optimisation procedure that is a component of the MAGIC suite (i.e., 226 MAGICOPT), generally used for regional applications. The minimum required site-measured 227 variables for calibration are: surface water concentrations for the major ions and soil 228 exchangeable fractions for base cations: calcium (Ca^{2+}), magnesium (Mg^{2+}), sodium (Na^{+}) 229 and potassium (K^+). The MAGIC suite also includes an iteration routine for the determination 230 of target loads. In this study, the deposition path was optimised between 2010 (Gothenburg 231 Protocol) and 2020 (the implementation year) to ensure the ANC-limit was achieved by 2050.

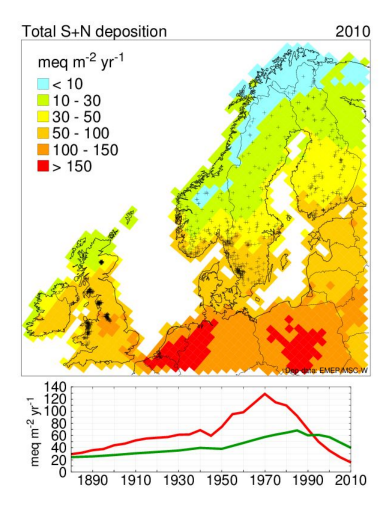
232

233 In the current study, the application of MAGIC (by national experts) across the study lakes followed a common (prescribed) procedure as described in Posch et al.³⁸; for further details 234 235 on the application and calibration of MAGIC see, for example, Aherne et al.⁴⁴ In brief, all 236 catchments were represented by one soil compartment receiving deposition and releasing 237 discharge to the lake compartment. The soil compartment represented the aggregated horizons 238 of the catchment soils (mineral and organic), with exchangeable base cation data taken from 239 national and focused surveys.³⁸ Simulations were carried out using an annual time-step, with a 240 number of simplifying assumptions applied consistently across all study lakes. Discharge 241 volume and flow routing within the catchments were not varied; discharge was described 242 using long-term means with 100% routed to the lake. Detailed process-oriented N dynamics 243 were not modelled, i.e., the coefficient for in-lake N retention was set to a similar value for all

244 lakes³⁶, and terrestrial N retention was set to match observed lake concentrations. To account 245 for uncertainty in a number of the fixed parameters (lake retention, soil depth, soil bulk 246 density, cation exchange capacity, etc.), a 'fuzzy' optimisation method was employed. Ten 247 calibrations were carried out for each study lake using MAGICOPT; for each simulation the 248 fixed parameters were randomly varied within specified uncertainty bands ($\pm 10\%$). Uptake of 249 base cations from forested catchments were modelled using a simplified regional sequence, 250 based on a regional average planting date, constant nutrient concentrations (from literature), 251 and species composition and coverage from national forest inventories (for further details see, e.g., Aherne et al.^{52,55} for Finland and Moldan et al.⁴⁶ for Sweden). 252 253 254 In the current study, MAGIC was calibrated to 848 lakes (see Figure 2) across Finland 255 (FI=163), Norway (NO=131), Sweden (SE=234), and the United Kingdom (UK=320),

spanning a range in deposition (S plus N) from <10 meq m⁻² yr⁻¹ in northern Sweden and 256 257 Finland to 150 meq m⁻² yr⁻¹ in the South Pennines, UK (Figure 2). Historic and future surface 258 water chemistry for each lake were simulated for the period 1880-2100 under modelled anthropogenic S and N deposition⁵⁶ from the EMEP model.⁵⁷ Future lake chemical recovery 259 260 under the Gothenburg Protocol (which came into force in 2010) was evaluated using charge 261 balance ANC (defined as the difference between the sum of the concentrations of base cations 262 and strong acid anions); lake ANC is an established acidification indicator, as it is strongly indicative of biological recovery.^{58,59} In addition, ANC is the most widely used chemical 263 264 criterion in critical load calculations for surface waters.⁶⁰ Target loads were estimated for the 265 target year 2050, with the implementation year 2020, i.e., the year in which deposition reductions beyond the Gothenburg Protocol (year 2010) are fully implemented to ensure 266 267 attainment of the critical chemical criterion by the target year.³⁸ While dynamic modelling 268 was carried out on every study site (n = 848), target loads were only calculated for each 269 surface water that did not meet the specified critical chemical criterion (ANC-limit) by 2050 270 under the Gothenburg Protocol, i.e., those lakes that still violated ANC-limits or with a 271 recovery delay (Note: lakes were the only study sites that still violated the ANC-limit). The 272 specified ANC-limit varied among countries (based on national management objectives); all 273 Swedish lakes had a fixed value of 20 meg m⁻³, similarly the UK surface waters (lakes and 274 streams) had a value of 20 meg m⁻³, except for a small number of naturally acidic sites that 275 had a limit of zero (n=21; 6.5%). Organic acids can act as strong acid anions reducing the acid 276 neutralizing (buffering) capacity of a lake to incoming acidity⁶¹; to accommodate this, Finland 277 and Norway used an organic acid adjusted ANC-limit⁶², which, for example, resulted in an 278 average ANC-limit of 14 meg m⁻³ for the Finnish lakes.

279



280

Figure 2: Location of the study sites used for dynamic modelling (n = 848; black crosses) overlaid upon modelled total sulphur (S) plus nitrogen (N) deposition in 2010 (meq m⁻² yr⁻¹) on the EMEP 50 km × 50 km grid (for further details on the EMEP model see Simpson et al.⁵⁷). The temporal development (1880–2010) of the total S (red) and N (green) deposition averaged over the 848 study sites is also shown.

286

287 Target load functions were estimated for each calibrated surface water that did not meet the

- 288 specified critical chemical criterion (ANC-limit) by 2050 under the Gothenburg Protocol, i.e.,
- every pair of N and S deposition that met the ANC-limit in 2050 under further (beyond
- 290 Gothenburg) emission reductions defined a target load function of acidity (TLF), similar to a
- 291 critical load function²¹ (see also Supporting Information for further details) for each study
- 292 lake (catchment). The piece-wise linear function in the (N, S) deposition-plane is delineated
- by the maximum target load of S, TLmaxS (for N deposition = 0) and the maximum target
- load of N, TLmaxN (for S deposition = 0).
- 295

296 4. Results and Discussion

Regional dynamic modelling results have been reported for individual countries.^{42,45,46,52,63} However, previous assessments primarily focused on scenario analyses, i.e., simulations to answer the question: 'what is the future chemical status of a surface water under various deposition scenarios?' In contrast, the current study addresses the inverse question: 'what deposition, called target load, is required to obtain a specified lake chemical status within a given time period (if feasible)?'.

303

Dynamic model simulations were carried out for 848 surface waters, but target load 304 305 calculations were only necessary for 173 lakes (Table 1). The simulated water chemistry for 306 the target year 2050 was predicted to be greater than (or equal to) the chemical criterion 307 (ANC-limit) for 675 surface waters. Target loads, i.e., loads below the respective critical 308 loads, were determined for 24 lakes in Finland, 56 in Norway, 14 in Sweden and 79 in the 309 United Kingdom. Of these 173 lakes, 46 were 'infeasible' (Table 1), i.e., even reducing 310 anthropogenic deposition to zero by 2020 would not result in an ANC greater than or equal to 311 the ANC-limit in 2050. In general, infeasible lakes occurred in < 3% of the study sites per country; however, in the United Kingdom, infeasible lakes occurred in $\sim 11\%$ (n = 35) of the 312 313 study sites likely reflecting their higher cumulative historic deposition (Figure 2). Neglecting 314 infeasible sites, 'true' target loads were determined for127 lakes (Table 1), 21 in Finland, 52 315 in Norway, 10 in Sweden and 44 in the United Kingdom; the highest proportion occurred in 316 Norway (40%) followed by Finland (13%).

317

The average TLmaxS (see Supporting Information) per country ranged from 7.5 meq m⁻² yr⁻¹ (Finland) to 38.9 5 meq m⁻² yr⁻¹ (United Kingdom). Note, for all study sites the maximum critical load of S (CLmaxS) was also computed as the steady-state solution of the dynamic model; compared with the average CLmaxS, the average TLmaxS was 53% lower in Finland, 40% in Norway, 20% in Sweden and 36% in the United Kingdom. Across all lakes, average TLmaxS (24.1 meq m⁻² yr⁻¹; n = 127) was 38% lower than the respective CLmaxS (39.1 meq m⁻² yr⁻¹; Table 1).

325

Table 1: Number of lakes in each country with dynamic model (DM) simulations, divided into the

327 number of lakes for which the critical load is sufficient to achieve the ANC-limit in 2050 (ANC-2050),

number of infeasible sites (INF) and 'true' target loads (TL). Also given are the averages of TLmaxS

329 and CLmaxS for lakes under 'TL'.

Country	DM	ANC-2050	INF	TL	TLmaxS*	S* CLmaxS	
				meq m ⁻² yr ⁻¹		$1^{-2} yr^{-1}$	
Finland (FI)	163	139	3	21	7.45	15.94	
Norway (NO)	131	75	4	52	19.17	31.94	

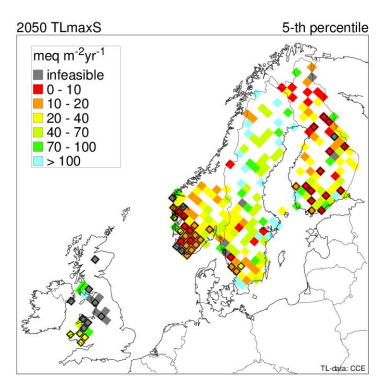
330	Sweden (SE)	234	220	4	10	19.91	25.02
	United Kingdom (UK)	320	241	35	44	38.87	61.72
-	Sum/Average	848	675	46	127	24.11	39.07

*See Supporting Information for further description of TLmaxS (and CLmaxS).

332

333 To provide greater regional coverage, target loads were mapped on the EMEP 50 km \times 50 km 334 grid (Figure 3) by setting TLmaxS to CLmaxS where TLmaxS > CLmaxS (since the critical load is already sufficient for non-violation of the ANC-limit by 2050). To account for all TLs 335 336 within a grid cell, the 5-th percentile of the cumulative distribution function for all target loads in that grid cell was mapped.²³ Overall, no clear pattern can be discerned in the mapped 337 338 target loads. In general, the critical load is sufficient for achieving non-violation of the ANC-339 limit in most areas; nevertheless 'true' target loads are concentrated in southern Norway and 340 Finland, and in northern Wales in the United Kingdom (Figure 3).

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Figure 3: Map of the 5-th percentile of the 2050 maximum target load of sulphur (TLmaxS) on the

345 Note: TLmaxS was set to the maximum critical load of S (CLmaxS) where TLmaxS > CLmaxS (i.e.,

the critical load is sufficient for non-violation of the ANC-limit by 2050). Grey-filled cells (label

347 'infeasible') denote grids containing at least one lake where the simulated ANC does not meet the

348 specified limit by 2050, even under zero deposition after 2020 (FI = 3, NO = 4, SE = 4, UK = 35; Table

349 1). Black diamonds frame grids with at least one 'true' TL, i.e., where a TL exists and is lower than the

- 350 CL (FI = 21, NO = 52, SE = 10, UK = 44; Table 1).
- 351

352 The key chemical variable of interest was ANC, as it is used as a chemical criterion linking

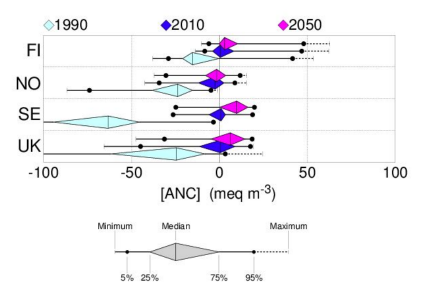
353 water chemistry to the biological (fish) status of the lakes; as such, time series of ANC were

354 simulated to illustrate the timing and rate of chemical changes during acidification and

355 recovery. The general pattern of predicted ANC in the study lakes is similar in the four

countries (Figure 4), driven by the deposition of S and N (Figure 2). The differences between

- 357 the regions were due to proximity to emission sources, acid sensitivity of regions, differences
- in land use and the selected lakes.
- 359



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Figure 4: Percentile statistics ('diamond plots') of simulated annual average lake acid neutralising
capacity (ANC) in 1990, 2010 and 2050 in Finland (FI), Norway (NO), Sweden (SE) and the United
Kingdom (UK). Data are only shown for lakes for which target loads were determined (FI = 24, NO =
56, SE = 14, UK = 79; see columns 'INF' plus 'TL' in Table 1).

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Implementation of the Gothenburg Protocol resulted in a significant increase in ANC from 366 367 1990 to 2010 (paired t-test, p < 0.001) in all four countries (average increase of 32.6 meq m⁻³; 368 Figure 4) and is predicted to significantly improve by 2050 albeit by a smaller amount 369 (average increase of 4.2 meg m⁻³; Figure 4). This is due to the fact that the deposition is kept 370 at the 2010 level after that year, and that only (slow) improvements in the soil and water conditions, e.g., replenishment of base cation pools, will raise the ANC.⁶⁴ Surface waters in 371 Sweden showed the greatest improvement in ANC between 1990 and 2010 (Figure 4) owing 372 to the concentration of sensitive lakes in south-western Sweden^{1,14} and the large reduction in 373 374 acidic deposition in that region (Figure 2). 375

376 Despite the predicted improvements in ANC, ultimately, we are concerned with the 'distance'

377 between the predicted chemical status of a lake and its desired status (note: the desired ANC-

378 limit is variable in all countries except in Sweden, see Figure 5a). A better characterisation of 379 the 'distance' of a lake's chemical status from the desired one is through the so-called ANC-380 deficit, i.e., the difference between simulated ANC (under the 2010 Gothenburg Protocol 381 deposition) and the (lake-specific) ANC-limit. The inverse cumulative distributions of the 382 ANC-deficit for the year 2050 were quite similar in the four countries up to the 80-th 383 percentile (Figure 5b); notably a maximum deficit around 40 meg m⁻³ was simulated for some 384 Norwegian lakes and 30 meq m⁻³ for some lakes in the United Kingdom. Implementation of target load depositions would ensure that all lakes reach their specified ANC-limit by 2050, 385 386 i.e., all ANC deficits reach zero by 2050.

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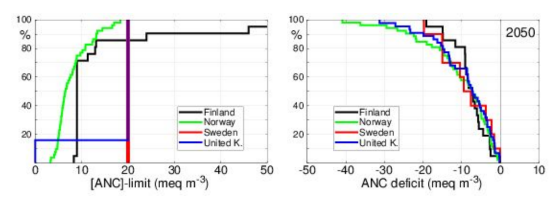




Figure 5: Left: Cumulative distribution functions of the ANC-limits for the lakes with target loads in
Finland (21), Norway (52), Sweden (10) and the United Kingdom (44). Right: Inverse cumulative
distribution functions of the 'ANC deficit' in 2050 for the same lakes in those countries. The ANC
deficit shows the difference between simulated ANC in 2050 under the 2010 Gothenburg Protocol (see
Figure 2) and the specified critical ANC-limit (left Figure).

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395 Dynamic modelling was carried out for 848 lake catchments in Finland, Norway, Sweden and 396 the United Kingdom. Given the large number of acid-sensitive lakes in each country, a larger 397 number would be desirable, but the chosen study lakes were deemed to be a representative 398 sample (as outlined in Materials and Methods) and (more importantly) have the inputs 399 required for dynamic modelling. It was predicted that 675 of these 848 lakes will achieve 400 their critical ANC-limit by the year 2050 under the Gothenburg Protocol, which leaves 173 401 lakes (20%), for which emission reductions beyond Gothenburg are required, if one wants to 402 achieve non-violation of the ANC-limit by 2050. However, for 46 of these lakes (~5% of the 403 total), even a reduction to zero deposition by 2020 would not be sufficient to achieve the ANC-limit by 2050. This does not mean that those lakes would never recover (chemically), 404 405 only that recovery would occur (maybe long) after 2050. 406 407 In the current study, model simulations have been conducted without consideration of future

climate change, as the primary objective was to support emissions reduction polices

409 (irrespective of climate perturbations). Nonetheless, several (regional) studies have been 410 conducted using MAGIC that explore the direct and indirect effects of climate change on lake 411 chemistry^{43,51,52}. Although the (indirect) effects can be great for individual lakes, the overall 412 effects on lake chemistry are not huge, considering all other (model) uncertainties.⁵¹ 413 Reductions of S and N deposition are the most important of determinants of future lake (acid) 414 status in European surface waters. 415 While target loads have been discussed and determined in Europe under the LRTAP 416 417 Convention, they have not been used explicitly to guide emission reduction policies. One 418 reason might be that it requires dynamic modelling – and thus more input data and expertise 419 to determine target loads - compared to critical loads that are 'easily determined' steady-state 420 quantities. However, lack of information on time needed for achieving the desired chemical 421 status under critical loads should ultimately encourage the determination of target loads to 422 provide policy advisors with guidance on the timing of ecosystem recovery. While 423 acidification is generally assumed to be 'solved' in Europe, there is growing recognition that 424 surface waters in some regions are still acidified²; the current assessment suggests that 425 emissions reductions beyond the Gothenburg Protocol are required to ensure surface water 426 recovery from acidification by 2050. 427 428 429 **Supporting Information** 430 Description of the Target Load Function (TLF) 431 432 Notes 433 The authors declare no competing financial interest. 434 435 Acknowledgements 436 M.P. thanks the Trust Fund for the effect-oriented activities under the Convention on Long-437 range Transboundary Air Pollution. This research was undertaken, in part, thanks to funding 438 to J.A. from an NSERC Discovery grant and the Irish EPA (2012-CCRP-MS.7 and 2016-439 CCRP-MS.43). M.F. thanks the Strategic Research Council at the Academy of Finland for 440 financial support (decision 312559). 441 442 References 443 (1) Garmo, Ø. A.; Skjelkvåle, B. L.; De Wit, H. A.; Colombo, L.; Curtis, C.; Fölster, J.; Hoffmann, A.; 444 Hruška, J.; Høgåsen, T.; Jeffries, D. S.; Keller, W. B.; Krám, P.; Majer, V.; Monteith, D. T.; 445 Paterson, A. M.; Rogora, M.; Rzychon, D.; Steingruber, S.; Stoddard, J. L.; Vuorenmaa, J.; 446 Worsztynowicz, A. Trends in surface water chemistry in acidified areas in Europe and North

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