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ENERGY, WATER, AND BROAD-SCALE GEOGRAPHIC PATTERNS OF SPECIES RICHNESS

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Abstract. It is often claimed that we do not understand the forces driving the global diversity gradient. However, an extensive literature suggests that contemporary climate constrains terrestrial taxonomic richness over broad geographic extents. Here, we review the empirical literature to examine the nature and form of the relationship between climate and richness. Our goals were to document the support for the climatically based energy hypothesis, and within the constraints imposed by correlative analyses, to evaluate two versions of the hypothesis: the productivity and ambient energy hypotheses. Focusing on studies extending over 800 km, we found that measures of energy, water, or water–energy balance explain spatial variation in richness better than other climatic and non-climatic variables in 82 of 85 cases. Even when considered individually and in isolation, water/energy variables explain on average over 60% of the variation in the richness of a wide range of plant and animal groups. Further, water variables usually represent the strongest predictors in the tropics, subtropics, and warm temperate zones, whereas energy variables (for animals) or water–energy variables (for plants) dominate in high latitudes. We conclude that the interaction between water and energy, either directly or indirectly (via plant productivity), provides a strong explanation for globally extensive plant and animal diversity gradients, but for animals there also is a latitudinal shift in the relative importance of ambient energy vs. water moving from the poles to the equator. Although contemporary climate is not the only factor influencing species richness and may not explain the diversity pattern for all taxonomic groups, it is clear that understanding water–energy dynamics is critical to future biodiversity research. Analyses that do not include water–energy variables are missing a key component for explaining broad-scale patterns of diversity.

Key words: *diversity gradients; energy hypothesis; latitudinal gradient; productivity hypothesis; species–energy theory; species richness; water–energy dynamics.*

INTRODUCTION

Almost two centuries after the discovery of the "latitudinal gradient" in species richness, a widely ac-

cepted explanation continues to elude ecologists and biogeographers. A sure indicator of our muddle is that, despite the large number of hypotheses that have been proposed (Pianka 1966, Huston 1979, 1994, Rohde 1992, Rosenzweig 1995), new hypotheses continue to appear (Ritchie and Olff 1999, Colwell and Lees 2000, Dynesius and Jansson 2000), and the total now exceeds

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thirty. But is it true that we lack sufficient evidence to reach a consensus regarding at least the primary factors influencing broad-scale richness gradients, especially at the global scale? We believe that even if we have not conclusively identified the primary mechanism(s) underlying global patterns of richness, it is at least possible to narrow substantially the list of most plausible explanations (see e.g., Rahbek and Graves 2001, Willig et al. 2003).

The energy hypothesis, a climatically based hypothesis that claims that energy availability generates and maintains richness gradients, is a likely candidate for a short list of explanations. The idea that climate's control of energy drives the global richness gradient dates from the beginning of biogeography (von Humboldt 1808) and has generated an extensive literature quantifying the relationship between species richness and climatic variables. Ten years ago, Wright et al. (1993) reviewed this literature and found broadly based support for the energy hypothesis. However, many additional studies of richness gradients have been conducted since then. Moreover, Wright et al. (1993) did not distinguish individual components of climate or consider the possibility that several mechanisms may underlie climate-richness relationships. In this paper, we return to this literature to address four related questions: (1) What is the scope of the empirical evidence that climate represents a primary explanation for richness gradients? (2) How much of the variation in species richness across broad scales can be accounted for by climate? (3) What specific aspects of climate best explain richness? and (4) How might two possible versions of the energy hypothesis contribute to richness gradients? The final question was not addressed by Wright et al. (1993), but is relevant because the energy hypothesis actually subsumes two driving mechanisms (Turner et al. 1987, 1996, Currie 1991, Hawkins et al. 2003).

One version of the hypothesis, as developed by Hutchinson (1959), Connell and Orias (1964), Brown (1981), and Wright (1983), proposes that energy constrains richness via trophic cascades. For example, Wright (1983) argued that at the base of the global food web, plant richness is limited primarily by solar energy and water availability (i.e., water-energy dynamics). Herbivore richness in turn is limited by the net primary production of plants, predator richness is limited by the secondary production of herbivores, and so on up the food chain. In this view, limits on richness are set by the energy flowing through food webs rather than the total energy entering a geographic area. This is often referred to as the productivity hypothesis and has received considerable attention (Wright et al. 1993, Huston 1994, Mittelbach et al. 2001).

The second hypothesis can be traced back to von Humboldt (1808) and is founded on the physiological requirements of organisms vis-à-vis ambient energy inputs rather than food availability. For example, von

Humboldt argued that plant richness declines at higher latitudes because many species are frost-intolerant and cannot survive cold temperate-zone winters. These cold winters are a direct result of seasonal declines in ambient energy as one moves away from the equator. With respect to animals, Turner et al. (1987) argued that sunshine and summer temperatures constrain butterfly richness in Great Britain because adult activity levels depend on ambient temperature and on basking in direct sunlight. Similarly, Currie (1991) hypothesized that thermoregulatory needs explain why the species richness of terrestrial North American vertebrates is more strongly correlated with annual potential evapotranspiration (PET, a measure of ambient or atmospheric energy) than with annual actual evapotranspiration (AET, a measure of water-energy balance closely associated with plant productivity). Whether or not the explanations proposed by von Humboldt (1808), Turner et al. (1987), and Currie (1991) are correct in detail, they represent an alternative mechanism to that represented by the productivity hypothesis. Of course, both physiological and trophic factors may underlie associations between climate and richness, and an additional focus of our analysis is to evaluate where on the globe and for which taxa pure energy variables or water-energy variables best predict diversity gradients.

METHODS

A database of studies of broad-scale richness gradients was compiled by a National Center for Ecological Analysis and Synthesis (NCEAS) working group on species-energy theory. Computerized literature databases, journals, the database generated by Wright et al. (1993), and a productivity-richness database generated by a previous NCEAS working group (Mittelbach et al. 2001) were all searched. A study was included if we could extract coefficients of determination derived from any standard statistical technique for individual explanatory variables for variation in the taxonomic richness (usually species, but occasionally generic or familial) of any terrestrial plant or animal group, although we also included a few studies where pairs of variables were combined by authors. In a few cases, raw data could be extracted from the original paper. If so, we recalculated the models to verify the published results, transforming variables if appropriate and computing coefficients of determination if the original authors had not provided them. Because these data were reanalyzed, the coefficients of determination sometimes differ from those reported in the original publications.

It is clear that the factors influencing richness gradients are scale dependent, and thus, explanations may vary with the spatial scope of the analysis (extent) and with sample resolution (grain) (Willis and Whittaker 2002). Our focus here is on gradients extending over at least 800 km of linear distance. We considered this distance minimally sufficient to encompass a range of

climates over which richness may vary. We also examined possible influences of grain size by recording whether the richness data had been generated from range maps (map data) or by sampling flora or fauna (sample data). Maps are usually used for large-grained studies (resolution measured in thousands to hundreds of thousands of square kilometers), whereas fine-grained studies (resolution measured in square meters to hectares) use samples of the taxa actually present in plots.

We excluded from consideration papers focused entirely on area, latitude, altitude or “geometric constraints.” Clearly, larger areas generally support more species than smaller areas, *as long as the climates among the areas are not too different*. The importance of climatic consistency among areas becomes obvious if we ask “Which area supports the most plant and animal species, Costa Rica (~51 000 km²), the Sahara Desert (~9 065 000 km²), or Antarctica (~13 340 000 km²)?” If this comparison seems absurd, it is only because most ecologists accept that climate must be taken into consideration when comparing areas with widely different climatic regimes. As such, analyses that exclude climatic variables when examining species–area relationships across large extents are likely to confound the effects of both variables and cannot be unambiguously interpreted. Second, virtually all workers realize that statistical relationships between richness and latitude or altitude provide no information about underlying driving factors, so studies focusing on these variables were excluded. Finally, we excluded papers focused on geometric constraints on diversity; operationally, because, like studies of latitude or area, correlating diversity with points on the earth’s surface without also including other variables makes it impossible to assess the extent that the correlations are due to covariation with spatially patterned environmental factors; more fundamentally, it has been argued that this idea is based on a flawed range concept and cannot explain diversity gradients in either real or null worlds (Hawkins and Diniz-Filho 2002).

RESULTS

We located 85 data sets: 22 for plants, 44 for vertebrates, and 19 for invertebrates (Table 1). Because of fundamental biological differences among these three groups, we evaluate each separately. However, across all groups, we found that the single best predictor variable explained, on average, 63.4% of the variance in richness (range 22–98%). The amount of variance explained does not depend on whether the data are map-vs. sample-based (ANOVA, $F_{1,83} = 0.004$, $P = 0.95$, map mean = $63.3 \pm 2.4\%$, sample mean = $63.6 \pm 3.4\%$ [means ± 1 SE]) or on which group is considered (ANOVA, $F_{2,82} = 1.44$, $P = 0.24$, plants mean = $68.9 \pm 3.8\%$, vertebrates mean = $61.2 \pm 2.7\%$, and invertebrates mean = $62.1 \pm 4.1\%$). We also found that climatic variables (whether direct or indirect via plant

productivity) were the strongest predictors of richness in 83 of the 85 cases, offering widespread support for the hypothesis that climate in general has a major influence on diversity gradients across large spatial extents. However, there also is clear taxonomically and geographically based heterogeneity in the data with respect to the specific climatic variables most strongly associated with richness, relevant to the question of whether water, energy, or their interaction may underlie diversity gradients.

Plant richness

In 20 of 21 cases, variables including water inputs emerge as the primary predictor of plant richness (Table 1), based on the r^2 values of each tested variable. The specific variables found reflect the variables included in each study, but all represent measures of water (rainfall and precipitation) or water–energy (actual evapotranspiration, productivity, net primary productivity, and net aboveground productivity). Further, water variables tend to be the best predictors when the geographic scope of the data is restricted to tropical and subtropical areas, whereas water–energy variables dominate when colder areas are also included. The single exception to the general result is for North American trees, where a reanalysis of Currie and Paquin’s (1987) data found that a temperature index explained two-thirds of the variance (Table 1; Allen et al. 2002). However, the analysis of Allen et al. (2002) was specifically designed to support a temperature-based hypothesis and did not include any water-related variables. Clearly, the data strongly implicate water availability as a key constraint on plant richness, particularly in warm areas where energy is abundant. In cold regions where energy inputs are lower and thus more likely to be limiting, energy interacts with water to explain richness gradients. It is notable that energy inputs alone never represent the primary explanatory variable for plant richness gradients, unless both water and water–energy variables are excluded from the analysis. Thus, a parsimonious interpretation of the plant studies is that plant richness is determined largely by water–energy dynamics, although pure water variables may be most strongly associated with diversity in warm areas.

Vertebrate richness

The relationships between water, energy, and richness show a wider range of variation for vertebrates than for plants (Table 1). In 24 cases, water or water–energy (measured either directly or indirectly as plant productivity) explained the most variance in richness, whereas in 17 cases, direct energy measures dominate. However, there are taxonomic and geographical patterns in the results. First, seven of the latter cases involve reptiles/lizards, suggesting that the richness of at least some reptile groups is most strongly constrained by ambient energy inputs, wherever they occur.

TABLE 1. Primary explanatory variables in analyses of broad-scale species richness patterns (i.e., extent > 800 km) of plants, vertebrates, and invertebrates; other classes of variables included in each study are also listed.

| Taxonomic group | Geographic area | Data type† | Primary variable‡ | Other variables§ | r ² | Reference |
|---------------------|-------------------------|------------|--------------------|------------------|----------------|---|
| Plants | | | | | | |
| Plants | global | samples | productivity | E,W,WE,T,A,O | 0.28 | Scheiner and Rey-Benayas (1994) |
| Plants | global (islands) | maps | AET | A | 0.76 | Wright (1983) |
| Plants | California | maps | precipitation | E,T,A,O | 0.52 | Richerson and Lum (1980) |
| Angiosperm families | global | maps | PET + WD | O | 0.84 | Francis and Currie (2003) |
| Woody plants | global | samples | rainfall | T,O | 0.74 | Gentry (1988) |
| Woody plants | N. Neotropics | samples | rainfall | ... | 0.86 | Gentry (1982) |
| Woody plants | N. Neotropics | samples | rainfall | T,O | 0.53 | Clinebell et al. (1995) |
| Woody plants | S. Africa | maps | rainfall | E,WE,T,O | 0.60 | O'Brien (1993) |
| Woody plants | Australia | samples | productivity | ... | 0.98 | Specht and Specht (1994) |
| Trees | N. temperate zone | samples | AET | A,O | 0.67 | Francis and Currie (1998) |
| Trees | N. America | maps | AET | E,W,T,O | 0.62 | Currie and Paquin (1987) |
| Trees | N. America | maps | NPP (modeled) | E,T,O | 0.91 | Adams and Woodward (1989) |
| Trees | Europe | maps | NPP (modeled) | E,T,O | 0.84 | Adams and Woodward (1989) |
| Trees | E. Asia | maps | NPP (modeled) | E,T,O | 0.98 | Adams and Woodward (1989) |
| Trees | S. America | samples | rainfall | ... | 0.65 | Kay et al. (1997) |
| Trees | Madagascar | samples | rainfall | ... | 0.50 | Ganzhorn et al. (1997) |
| Trees | N. America | maps | temperature | ... | 0.66 | Allen et al. (2002) |
| Bog plants | northeastern N. America | samples | rainfall | E,A,O | 0.60 | Glaser (1992) |
| Herbs/shrubs | Chile | samples | rainfall | ... | 0.70 | Meserve and Glanz (1978) |
| Pteridophytes | Bolivia | samples | precipitation | E,T,A,O | 0.64 | Kessler (2001) |
| Cacti | Argentina | maps | summer rainfall | E,WE,O | 0.50 | Mourelle and Ezcurra (1996) |
| Grasses | Namibia | samples | rainfall | ... | 0.77 | Schulze et al. (1996) |
| Vertebrates | | | | | | |
| Vertebrates | Afrotropics | maps | NPP (modeled) | T,O | 0.73 | Balmford et al. (2001) |
| Vertebrates | N. America | maps | PET | W,WE,T,O | 0.92 | Currie (1991) |
| Birds | Global (islands) | maps | NPP (modeled) | A | 0.80 | Wright (1983) |
| Birds | Global (continents) | maps | AET | E,W,T,O | 0.72 | Hawkins et al. (2003) |
| Birds | India | maps | rainfall | ... | 0.51 | Pearson and Carroll (1998) |
| Birds | S. Africa | maps | precipitation | E,WE,O | 0.67 | van Rensburg et al. (2002) |
| Birds | Australia | maps | rainfall | E,O | 0.48 | Pianka and Schall (1981) |
| Birds | USA + southern Canada | samples | NDVI | T | 0.61 | Hurlbert and Haskell (2003) |
| Birds | USSR | maps | temperature | W | 0.89 | Terent'ev (1963) |
| Birds | Norway | samples | May temperature | ... | 0.78 | Heggberget (1987) |
| Birds (summer) | UK | maps | summer temperature | W,T,O | 0.34 | Lennon et al. (2000) |
| Birds (winter) | UK | maps | summer temperature | W,T,O | 0.59 | Lennon et al. (2000) |
| Passerine birds | Argentina | maps | temperature | W,T,O | 0.57 | Rabinovich and Rapoport (1975) |
| Amphibians | Europe | maps | AET | E,W,P,T,O | 0.61 | M. A. Rodriguez, J. A. Belmontes, and B. A. Hawkins (<i>unpublished data</i>) |

TABLE 1. Continued.

| Taxonomic group | Geographic area | Data type† | Primary variable‡ | Other variables§ | r^2 | Reference |
|---------------------------|--------------------|------------|----------------------------|------------------|-------|---|
| Amphibians | Iberia | maps | precipitation | E,O | 0.22 | Schall and Pianka (1977) |
| Amphibians | N. America | maps | temperature | ... | 0.69 | Allen et al. (2002) |
| Amphibians | USSR | maps | temperature | W | 0.46 | Terent'ev (1963) |
| Frogs | USA | maps | rainfall | E,T,O | 0.64 | Schall and Pianka (1978) |
| Frogs | Australia | maps | rainfall | E,O | 0.53 | Pianka and Schall (1981) |
| Mammals | N. Neotropics | samples | rainfall | T,A,O | 0.48 | Medellin (1994) |
| Mammals | Chile | samples | precipitation | O | 0.90 | Meserve and Glanz (1978) |
| Mammals | USSR | maps | temperature | W | 0.89 | Terent'ev (1963) |
| Mammals | N. America | maps | temperature | W,WE,T | 0.57 | Badgley and Fox (2000) |
| Mammals | S. Africa | maps | variability in temperature | W,WE,T,O | 0.69 | Andrews and O'Brien (2000) |
| Mammals | N. Palearctic | maps | percentage of forest | E,O | 0.79 | Danell et al. (1996) |
| Mammals | N. Nearctic | maps | forest fragmentation | E,O | 0.45 | Danell et al. (1996) |
| Marsupials | Australia | maps | rainfall | E,O | 0.54 | Pianka and Schall (1981) |
| Primates | S. America | samples | rainfall | P,O | 0.37 | Kay et al. (1997) |
| Primates | S. America | samples | rainfall | ... | 0.67 | Reed and Fleagle (1995) |
| Primates | Africa | samples | rainfall | ... | 0.75 | Reed and Fleagle (1995) |
| Primates | Madagascar | samples | rainfall | ... | 0.70 | Reed and Fleagle (1995) |
| Lemurs | Madagascar | samples | rainfall | O | 0.46 | Ganzhorn et al. (1997) |
| Rodents | southwestern USA | samples | precipitation | O | 0.44 | Brown (1973) |
| Reptiles | Europe | maps | PET | W,WE,P,T,O | 0.71 | M. A. Rodriguez, J. A. Belmontes, and B. A. Hawkins (<i>unpublished data</i>) |
| Reptiles | USSR | maps | temperature | W | 0.90 | Terent'ev (1963) |
| Reptiles | Iberia | maps | hours of sunshine | W,O | 0.36 | Schall and Pianka (1977) |
| Snakes | Australia | maps | rainfall | E,O | 0.57 | Pianka and Schall (1981) |
| Snakes | USA | maps | rainfall | E,T,O | 0.32 | Schall and Pianka (1978) |
| Turtles | Australia | maps | rainfall | E,O | 0.77 | Pianka and Schall (1981) |
| Turtles | USA | maps | rainfall | E,T,O | 0.53 | Schall and Pianka (1978) |
| Lizards | USA | maps | hours of sunshine | W,T,O | 0.67 | Schall and Pianka (1978) |
| Lizards | western N. America | samples | July temperature | W,T,O | 0.88 | Pianka (1967) |
| Lizards | western USA | samples | Jan temperature | W,T,O | 0.32 | Scheibe (1987) |
| Lizards | Australia | maps | temperature | W,O | 0.44 | Pianka and Schall (1981) |
| Invertebrates | | | | | | |
| Tiger beetles | N. America | maps | PET | W,WE,O | 0.87 | Kerr and Currie (1999) |
| Tiger beetles | India | maps | rainfall | ... | 0.58 | Pearson and Carroll (1998) |
| Tiger beetles | Australia | maps | rainfall | ... | 0.25 | Pearson and Carroll (1998) |
| <i>Epicauta</i> (beetles) | N. America | maps | PET | W,WE,T,O | 0.82 | Kerr and Packer (1999) |
| Dung beetles | France | maps | temperature | W,T,O | 0.36 | Lobo et al. (2002) |
| Butterflies | W. Palearctic | maps | AET | E,W,T,O | 0.79 | B. A. Hawkins and E. E. Porter (<i>unpublished data</i>) |

TABLE 1. Continued.

| Taxonomic group | Geographic area | Data type† | Primary variable‡ | Other variables§ | r^2 | Reference |
|--------------------|------------------|------------|--------------------------------------|------------------|-------|--|
| Butterflies | Afrotropics | maps | AET | E,W,T,O | 0.69 | B. A. Hawkins and E. E. Porter (<i>unpublished data</i>) |
| Butterflies | Australia | maps | AET | E,W,T,O | 0.62 | B. A. Hawkins and E. E. Porter (<i>unpublished data</i>) |
| Butterflies | Australia | maps | summer rainfall | E,T,O | 0.64 | Dingle et al. (2000) |
| Butterflies | UK | maps | summer sunshine + summer temperature | ... | 0.79 | Turner et al. (1987) |
| Butterflies | N. America | maps | PET | W,WE,T,O | 0.69 | B. A. Hawkins and E. E. Porter (<i>unpublished data</i>) |
| Butterflies | Canada | samples | landcover diversity | E,W,WE,T | 0.74 | Kerr et al. (2001) |
| Moths | UK | maps | summer sunshine + temperature | ... | 0.72 | Turner et al. (1987) |
| Forest Lepidoptera | Canada | maps | PET | WE,T,O | 0.69 | Kerr et al. (1998) |
| Termites | Africa | maps | NPP | ... | 0.39 | Eggleton et al. (1994) |
| Termites | Neotropics | maps | NPP | ... | 0.41 | Eggleton et al. (1994) |
| Termites | Indo-Malaysia | maps | NPP | ... | 0.46 | Eggleton et al. (1994) |
| Ants | W. Hemisphere | maps | NAP (modeled) | ... | 0.58 | Kaspari et al. (2000) |
| Ants | southwestern USA | samples | rainfall | E,O | 0.70 | Davidson (1977) |

Note: The coefficients of determination (r^2) refer to the explanatory power of the primary variable only.

† Maps = richness generated using range maps, and samples = richness estimated from samples.

‡ Abbreviations are: AET = actual evapotranspiration; WD = water deficit; PET = potential evapotranspiration; NPP = net primary productivity; NDVI = normalized difference vegetation index; NAP = net aboveground productivity.

§ Abbreviations are: E = energy, W = water, WE = water–energy, P = productivity; T = topography, A = area, O = other, and ellipses (...) = none.

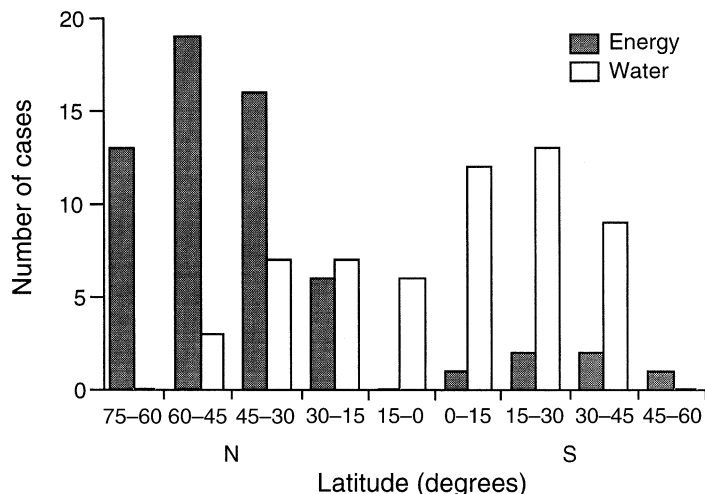
Second, eight of the energy cases not involving reptiles represent studies encompassing regions with widespread cold climates (North America, Norway, Great Britain, and the former Soviet Union). This is in contrast to studies identifying either water as the best predictor (in regions dominated by warm climates) or water–energy variables (in globally extensive data and one case in Europe). However, there are four apparent exceptions. The first involves passerine birds in Argentina (Rabinovich and Rapoport 1975). A major portion of Argentina is mild temperate, yet temperature emerged as the best predictor of bird richness. On the other hand, southern Argentina does include sub-Antarctic climatic areas, and it is likely that temperature is limiting in the far southern hemisphere, as it is in the far northern hemisphere. Another exception was found for the mammals of southern Africa, but in this case, the best climatic predictor was neither energy nor water, but a measure of seasonality (Andrews and O'Brien 2000). This represents the only case we found that identified seasonal variability as predicting richness patterns better than annual climatic variables, and as such, represents an exception to the more general patterns. A study

of herbivorous mammals in the northern boreal zone (Danell et al. 1996) also appears to be exceptional, as measures of forest size were the best predictors of richness in both the Palearctic and Nearctic (Table 1). However, based on their full analysis, they concluded that mammal richness was probably temperature limited, consistent with other studies of mammals in the northern temperate zone (Table 1). Thus, despite some possible exceptions, the data for most vertebrates indicate that water–energy variables best account for global-scale patterns, whereas in warm climates, water or water–energy is most strongly associated with richness, and in cold climates ambient energy inputs become critical.

Invertebrate richness

Although there are fewer data, invertebrates show a pattern very similar to that for vertebrates (Table 1). Water–energy or water variables are the primary predictors, except in some studies encompassing colder regions in the northern temperate zone (North America, Great Britain, and Canada). The case of dung beetles in France might represent an exception, in that they are

FIG. 1. The latitudinal distribution of explanatory variables in 38 analyses of animal richness gradients in which pure energy or pure water variables best explained richness. Each case was assigned to one or more latitudinal bands depending on its geographical scope, and the primary explanatory variable for the study was scored in each band. All scores were then summed within each band. For example, in the 13 cases where at least part of the richness data occurred between 75° N and 60° N, all found energy as the best predictor, whereas none found water to be the best predictor.



farther south and temperature is still the primary predictor (Lobo et al. 2002), but we will return to this case later. A study of Canadian butterflies (Kerr et al. 2001) represents a case in which climate was not the best predictor, although it could be argued that land-cover diversity is itself under climatic control. Nevertheless, the geographic distribution of studies of invertebrates is, overall, consistent with what has been found in vertebrates; energy has a strong relationship with richness in cool climates, whereas water by itself or in conjunction with energy is critical in warmer climates.

Latitude and constraints on diversity

As indicated in the previous two sections, studies focused on vertebrates or invertebrates suggest a shift in the primary limiting factor from energy to water as one moves from the poles to the equator. To explore this further, we selected the 38 cases for which animal richness patterns were best described by pure energy (PET or temperature) or pure water (rainfall or precipitation), which allowed us to isolate the separate components of water–energy. We then divided the world into nine latitudinal bands of 15° each, and scored which bands each study encompassed. We then summed all of the studies within each band with respect to the best explanatory variable and plotted the totals against latitude. Although we cannot analyze the resulting pattern statistically because most studies contributed data to multiple bands, the pattern (Fig. 1) reinforces the conclusion that the primary constraint on animal richness varies latitudinally. In high latitudes of the northern hemisphere, energy is most often the primary explanatory variable, with increasing numbers of cases in which water is the strongest predictor as we move south. In the southern hemisphere, water is by far the dominant explanatory variable, except in cases involving reptiles and Argentinean birds. It also appears that the latitudinal transition in the northern

hemisphere from energy to water as the primary predictor is gradual, extending over 45° of latitude, and there a strong asymmetry between the northern and southern hemispheres as a whole (Fig. 1). We will return to these issues in the *Discussion*.

DISCUSSION

Our synthesis extends the earlier work of Wright et al. (1993) and provides basic answers to the questions raised in the *Introduction*. Climate is typically a strong descriptor of broad-scale richness patterns, and the single best climatic predictor on average explains over 60% of the variance. Also, both water and energy inputs play a prominent role in richness patterns of a wide range of taxonomic groups comprising both plants and animals, supporting the hypothesis that “water–energy dynamics” provides a potent explanation for the diversity gradient across the entire planet (see e.g., O’Brien 1998). Finally, the geographic pattern found among the studies also suggests that the relative importance of the two components of water–energy dynamics shifts latitudinally, such that energy places strong constraints on plant and animal richness in the far north (and probably the far south) where energy inputs are low, whereas water constrains richness in areas with high energy inputs. Thus, a major conclusion is that, if water drives plant production in warm climates as is widely assumed, both the ambient energy and productivity versions of the energy hypothesis find support, depending where in the world the study is focused and on overall water–energy inputs. However, although the data indicate a shift in the primary constraint on richness, the semi-quantitative method we used to compare across latitudes (Fig. 1) is not clear about where on the planet this shift may occur, or if the transition is gradual or abrupt. This is particularly true since curvilinear relationships between energy variables and richness are commonly found in analyses focused on northern latitudes. To examine this in more

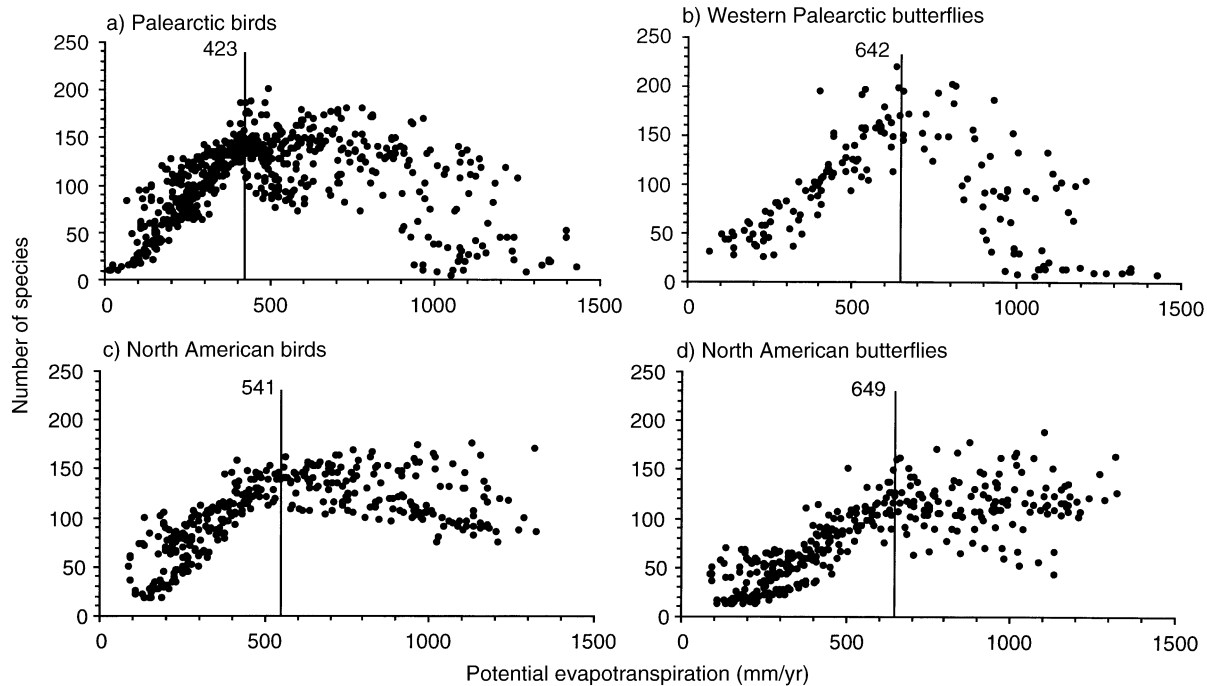


FIG. 2. Relationships between potential evapotranspiration (PET) and species richness for terrestrial birds (a, c) and butterflies (b, d) in the Palearctic (a, b) and Nearctic (c, d). The vertical lines represent breakpoints identified by split-line regression at which the relationships shift from being positive to either negative (in the Palearctic) or null (in the Nearctic). The regressions were performed using JMP 4.0 (SAS Institute 2000).

detail, we followed Kerr and Packer (1997) in using split-line regression to determine the nature of the relationship between energy and animal richness gradients in the northern hemisphere, where cold terrestrial climates are most widespread and, hence, where energy inputs are often low.

Currie's (1991) finding that vertebrate richness in North America was best described by PET was based on a curvilinear regression model. Kerr and Packer (1997) reanalyzed the mammal data and confirmed that richness was strongly positively associated with potential evapotranspiration above 45° – 48° N (near the USA–Canadian border), but not below this latitude. They proposed this boundary as the threshold below which ambient energy no longer limits mammal richness. To test the generality of this result, we examined the relationship between species richness and potential evapotranspiration using the terrestrial birds of North America and the Palearctic (Hawkins et al. 2003) and the butterflies of North America and the western Palearctic (B. A. Hawkins and E. E. Porter, unpublished data). Richness of both taxa was estimated in equal-area grid cells of 48 400 km² using range maps, and PET was estimated from a global database (available online).¹³

Consistent with other studies in the northern temperate zone (e.g., Currie 1991), both butterfly and bird

richness are strongly correlated with PET at low energy levels, but show either no relationship (Nearctic) or a negative relationship (Palearctic) at higher energy levels (Fig. 2). Split-line regressions further indicate that bird richness is primarily energy-limited only in Canada, the British Isles, Scandinavia, and across northern European Russia and Siberia (Fig. 3). Butterflies appear to be more sensitive to low energy than birds, showing strong positive relationships with PET across Canada and the extreme northern USA, and in most of Europe (Fig. 3). Although the abruptness of the shift from strong to weak associations of PET and richness is arguable, south of these lines it is probable that water replaces energy as the primary constraint (Table 1, Fig. 1). Whether these energy thresholds apply to other vertebrate and invertebrate groups remains to be determined, although the threshold for North American mammals (Kerr and Packer 1997) is very similar to that of North American butterflies, and French dung beetle richness is most strongly explained by temperature (Table 1), as would be predicted from the threshold found for European butterflies (Fig. 2). But based on the available evidence, it appears that energy is a strong predictor of animal diversity gradients in only a small part of the planet, and that over most of the earth the distribution of rainfall has a stronger influence on diversity gradients than temperature.

A final point about the geographic distribution of energy and water as predictors of diversity is that we

¹³ URL: <http://www.grid.unep.ch/data/grid/gnv183.html>

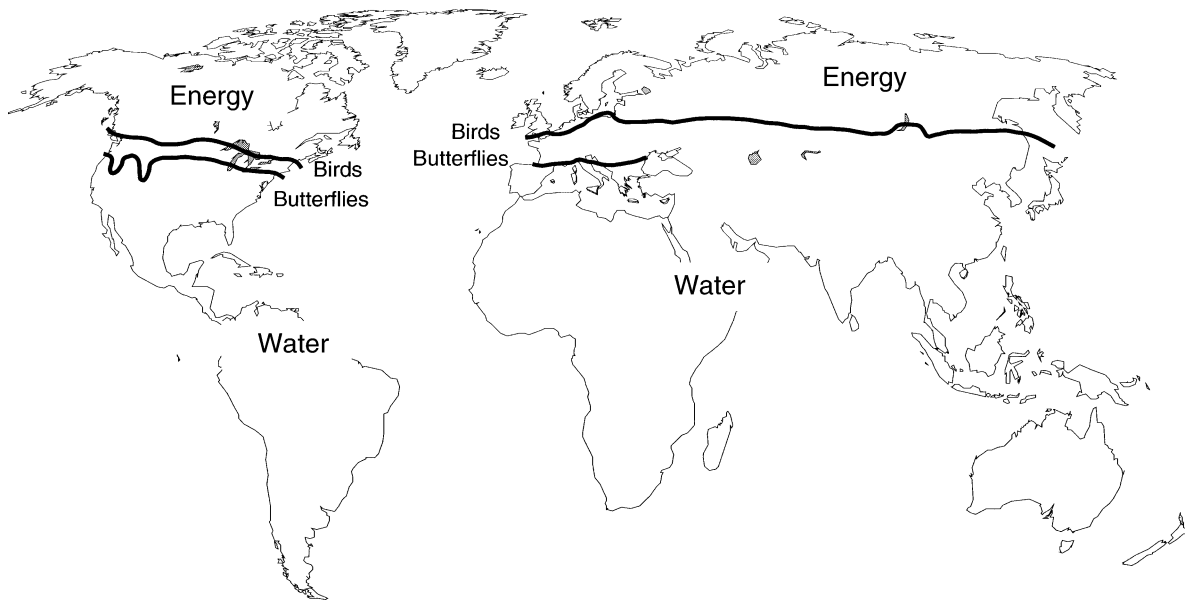


FIG. 3. A hypothesis for the geographic distribution of the limits to the species richness of animals, based on the analysis of butterflies and birds (see Fig. 2). The bold lines represent the geographical distribution of the breakpoints above which potential evapotranspiration and species richness are not positively associated. Across all latitudes, animal richness is constrained by the interaction of energy and water, but north of these lines energy is hypothesized to represent the limiting component of the interaction, whereas south of these lines water is assumed to be the key limiting component (see Fig. 1).

might expect the pattern to be roughly symmetrical around the equator, but it is not (Fig. 1). We consider it likely that the observed asymmetry reflects the fact that interior continental climates are primarily restricted to the northern hemisphere. In the southern hemisphere, climates in the temperate zone are generally much warmer, and even in the far south temperature fluctuations are more buffered by the oceans (outside of Antarctica there is no southern equivalent to Siberian winters), whereas precipitation can still be highly variable because of orographic effects. Thus, although low temperatures are likely to be critical in some parts of the southern hemisphere, such as southern Argentina and at high altitudes, the data suggest that water is more important across very broad geographic areas.

Reptiles, particularly lizards, are commonly found to be most closely associated with energy measures wherever they are studied. Because they are extreme solar ectotherms with a complex array of physiological and behavioral mechanisms for maintaining their body temperatures, it is believed that ambient temperature represents the most important climatic factor influencing reptiles (Heatwole 1976), a view supported by richness studies. The failure of the other major ectothermic vertebrate group, the amphibians, to show the same dependence on temperature at all levels is not surprising, given that most species require moisture to reproduce. Invertebrates are more puzzling. One potential explanation for why they do not follow the pattern found for reptiles is that their small body sizes create problems with desiccation in hot environments. Alter-

natively, both water and energy are critical to plants, and invertebrate richness may be linked to plant production, so the influence of water in warm environments is primarily indirect, operating via effects on food availability.

Whether energy and/or water influences both invertebrate and vertebrate richness gradients via direct physiological effects or via plant productivity is key to understanding the mechanism(s) underlying climatically driven richness gradients. However, in most cases, the alternative hypotheses are difficult to disentangle, particularly since they are not mutually exclusive. Plant productivity is closely associated with water–energy balance at the global scale (e.g., Rosenzweig 1968, Lieth 1975), and although new remote-sensing techniques now provide alternative measurements of net primary productivity (Goetz et al. 2000), in the past it has usually been modeled with climatic variables rather than measured directly (see also Huston 1999). Given these problems, it is difficult to decouple climatic vs. productivity influences on richness using multiple regression, the most common statistical technique. Consequently, care is necessary when attempting to determine whether it is more appropriate to refer to observed climate–richness links as supporting a physiologically based “water–energy dynamics” hypothesis or a tropically based “productivity” hypothesis, at least for animals. This complication does not apply to plants, as it is widely accepted that both water and energy are essential to plant physiological processes and thus have direct effects on plant abundance and diversity (Ste-

phenson 1990, O'Brien 1993). Given the broadly based evidence for the influence of water and energy on animal as well as plant richness, an important next step will be to determine the relative importance of direct and indirect climatic effects on geographic patterns of animals. The finding that pure energy variables usually describe animal diversity gradients best in the far northern latitudes suggests that direct physiological effects dominate in cold climates, but additional work is needed to confirm this interpretation.

A closely related issue is that a set of correlations among variables does not in itself show that they are linked by any causal mechanism, and there have been difficulties imagining mechanisms linking energy or water–energy to diversity gradients, especially if the theory is restricted to deterministic modeling (Rohde 1992). However, there are several models that predict such a direct relationship from either the turnover of species or the turnover of individuals. First, the classic species-turnover model that underlies the theory of island biogeography suggests that low energy (food) levels lead to low population sizes, and, consequently, higher extinction rates. This could lead to lower species numbers at dynamic equilibrium when energy levels are low (Hutchinson 1959, Turner 1992, Turner et al. 1996). Second, minimum viable population sizes may be lower in areas of high productivity (D. J. Currie et al., *unpublished manuscript*). This would also lead to greater species richness in warm, wet climates. Third, Hubbell's "neutral theory of biodiversity" (Hubbell 2001) uses a set of null assumptions about the turnover of individuals that predicts that species richness (via the "fundamental biodiversity number") will be directly related to the productivity and area of the metacommunity, as well as the speciation rate. Thus, there are three models which predict that population sizes, or the total number of individuals over all species, will influence the equilibrium number of species, and any environmental factor that influences population sizes across the appropriate spatial scale is therefore capable of influencing or controlling species richness. The productivity hypothesis proposes that the overriding factor is food supply. The ambient energy hypothesis assumes that growth and reproduction are greater at higher temperatures, and winter mortality also will be lower, leading to higher population sizes. This is obvious for plants and ectotherms, but is also true for mammals and birds, because higher overall temperatures allow endotherms to divert metabolic energy that might otherwise be spent on maintaining body temperature into growth and reproduction (Root 1988, Wright et al. 1993, Turner et al. 1996). Clearly, under very hot and arid conditions the availability of water per se will become the limiting factor.

Of course not all climatic variables, even those generally related to productivity, always have positive relationships with species richness over their entire range. Although curvilinear relationships may be found

across all spatial scales (Mittelbach et al. 2001), at large scales these are likely to occur because these variables do not in fact correlate with productivity over their entire range (e.g., O'Brien 1993, Scurlock and Olson 2002, Francis and Currie 2003). Alternatively, there are hierarchical controls on richness, illustrated by the relationships between PET and bird/butterfly richness in northern latitudes (see Fig. 2), where beyond a minimum threshold other factors become limiting.

Unquestionably, climate is strongly associated with richness gradients, but how well do climatic variables explain richness? In some cases, a single energy or water variable can by itself account for >90% of the variance in species richness (Table 1). But in others, the best predictor accounts for substantially less than half of the variance, raising the possibility that alternative, unmeasured factors may be better predictors. For example, it is widely assumed that the grain size and extent of the analysis has a strong influence on the processes driving richness. At small grain sizes and local extents a wide range of biotic and abiotic factors are important, whereas at large grain sizes and continental/global extents climatic and historical factors dominate (Whittaker et al. 2001, Willis and Whittaker 2002). There are also several examples showing how differences in grain size can influence the contributions of variables to regression models of richness (Rahbek and Graves 2001, Hurlbert and Haskell 2003). However, when we distinguished studies based on sampled data (i.e., richness measured over very local scales) vs. those based on range maps (i.e., richness estimated within large areas), we found no difference in the explanatory power of climate variables. Presumably, the range of climatic variation across the large extents considered here is sufficient to swamp the effects of any additional factors influencing richness, irrespective of the grain size. This issue will be further explored elsewhere. Even so, single climatic variables are sometimes insufficient to describe richness gradients very well, even over large spatial extents. In some of these cases, the inclusion of additional climatic variables substantially improved the predictive power of the models (see original papers listed in Table 1), but in others, most of the variance was not explained by any of the measured variables.

In closing, we note that our evaluation of the energy hypothesis should be viewed with some caveats. First, we do not claim that water–energy dynamics accounts for broad-scale richness variation in all taxonomic groups, nor that it is the sole factor driving richness of any single group. For example, the extent that the evolutionary history of different regions influences the global diversity gradient remains contentious (see e.g., Haydon et al. 1993, Latham and Ricklefs 1993, Schluter and Ricklefs 1993, McGlone 1996, Whittaker and Field 2000, Francis and Currie 2003, Hawkins et al. 2003), and we cannot exclude the possibility that biotic factors affect diversity at large grains and extents (e.g.,

Kaufman 1995). Even so, it is now clear that the link between water–energy and richness is widespread and generally strong, identifying it as a key component of future biodiversity research. Second, our conclusions are necessarily based on correlations rather than experimental evidence, making it difficult to tease apart direct and indirect effects of climate on richness and on underlying mechanisms. Thus, our interpretation must be tentative until mechanisms can be evaluated based on any mutually exclusive predictions each makes. Further, direct comparisons of energy to other hypotheses are challenging, owing to the varying natures of proposed explanations and to the difficulty in evaluating opposing hypotheses within a uniform framework. Finally, we have focused on terrestrial systems, although many of our arguments may also apply to marine and freshwater systems (e.g., Rex et al. 1993, Oberdorff et al. 1995, Roy et al. 1998, Rutherford et al. 1999). It is hoped that our synthesis will stimulate others to test the generality of these ideas in aquatic habitats.

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