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4 *Sedum* root foraging in layered green roof substrates

- 5 Peng Ji¹, Arne Sæbø², Virginia Stovin³, Hans Martin Hanslin^{2*}
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7 Addresses

- ⁸ ¹Agronomy College, Heilongjiang Bayi Agricultural University, Daqing, Heilongjiang,
- 9 People's Republic of China
- ²Department of Urban Greening and Environmental Engineering, Norwegian Institute
- 11 of Bioeconomy Research (Nibio), Ås, Norway
- ³Department of Civil and Structural Engineering, University of Sheffield, Sheffield, UK
- 13
- 14 * **Corresponding author**: hans.martin.hanslin@nibio.no
- 15

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- 29 Abstract
- 30

Background and aims Layered profiles of designed soils may provide long-term benefits
for green roofs, provided the vegetation can exploit resources in the different layers. We

33 aimed to quantify *Sedum* root foraging for water and nutrients in designed soils of

34 different texture and layering.

Methods In a controlled pot experiment we quantified the root foraging ability of the species *Sedum album* (L.) and *S. rupestre* (L.) in response to substrate structure (fine, coarse, layered or mixed), vertical fertiliser placement (top or bottom half of pot) and watering (5, 10 or 20 mm week⁻¹).

39 *Results* Water availability was the main driver of plant growth, followed by substrate 40 structure, while fertiliser placement only had marginal effects on plant growth. Root 41 foraging ability was low to moderate, as also reflected in the low proportion of biomass 42 allocated to roots (5-13%). Increased watering reduced the proportion of root length and 43 root biomass in deeper layers.

Conclusions Both *S. album* and *S. rupestre* had a low ability to exploit water and nutrients
by precise root foraging in substrates of different texture and layering. Allocation of
biomass to roots was low and showed limited flexibility even under water-deficient
conditions.

49 **Keywords:** green roof; Sedum; vegetation; root foraging; substrate texture and layering

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51 Introduction

52 Stormwater management through retention and detention on green roofs can be targeted 53 through the combinations of vegetation and soils used in the roof construction. The soils 54 are highly designed, usually lightweight and porous, to meet specific criteria for longterm functions. The role of the vegetation is to evaporate the stored water between rainfall 55 56 events and this is the limiting factor for stormwater management by green roofs in many 57 climates (Johannessen et al. 2017). While standard Sedum-based extensive green roofs 58 often function well across large climate gradients (Johannessen et al. 2017), vegetation 59 with higher water use or higher resistance and resilience to specific environmental conditions is sought to improve green roof functions, multifunctionality and stormwater 60 61 retention. Unfortunately, the use of non-succulent vegetation often entails a risk of 62 mortality and failure due to drought episodes (Johannessen et al. 2017; Monterusso et al. 2005; Nagase and Dunnett 2010). Therefore, further investigations of how green roofs 63 with *Sedum* species can be designed, could be useful to increase the role of green roofs 64 in stormwater management for the drier and wetter ends of the humidity gradient. 65

66

In coastal climates, *Sedum* may suffer winter damage as both shoots and roots are sensitive to prolonged wet conditions. One solution may be to use a coarse substrate on top to reduce moisture around shoots and a layered structure with a finer substrate deeper in the profile that is actually able to retain some water. Layered configurations may also be of wider interest, as high substrate temperature is a considerable problem for roof vegetation under dry Mediterranean conditions (Savi et al. 2016), but can be manipulated by substrate depths (Reyes et al. 2016) and to some extent by substrate composition 74 (Sandoval et al. 2017). Further, roots are less frost-tolerant than shoots and hence benefit 75 from substrates which they can forage into depths which are better frost insulated (Boivin 76 et al. 2001) and layered structures may better handle both water amounts and 77 contaminants (Wang et al. 2017). The feasibility of layered configurations is likely to 78 depend on the root foraging patterns of the vegetation and whether they are able to exploit 79 resources in vertical substrate layers. Despite their importance on green roofs, very little 80 is known about Sedum root systems and how the roots interact with the substrate and 81 environmental conditions to affect plant performance and green roof functions. A better 82 understanding of root foraging capacity and root growth patterns and knowledge of how 83 to manipulate these are steps towards more reliable *Sedum* based green roofs under contrasting climatic conditions. 84

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Plant root growth is governed by a set of plastic traits including branching patterns, root
diameter, specific root length and rooting depth, enabling roots to forage for resources
like water and nutrients (Hodge 2009). Root foraging is resource-demanding, so there is
clearly a trade-off with other plant functions and a link between foraging strategy, fitness
components and evolution (Jansen et al. 2009; Kembel and Cahill 2005; Weiser et al.
2016).

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Ecological limits to plastic responses like root foraging are expected when abiotic factors
have strong effects on plant fitness (Valladares et al. 2007). Stress-tolerant vegetation that
typically inhabit soils of small volume and low water-holding capacity, where abiotic
conditions including drought are of overriding importance, often have a low ability for

97 precise root foraging (Grime 2007; Grime and Mackey 2002) and may depend more on reducing water loss to survive adverse periods. Succulent leaves and different degrees of 98 99 crassulacean acid metabolism (CAM) are parts of an adaptive suite of traits under such conditions. Succulents often also have a low allocation of biomass to roots (Poorter et al. 100 101 2012; von Willert et al. 1991), shallow root systems with wide lateral spread (Schenk and 102 Jackson 2002) and rely on opportunistic water acquisition during wet periods and storage 103 between rain events. Roots of some succulents are also found to rapidly restore function 104 on rewetting and to have a low loss of water to drying soil (Nobel and Huang 1992; Nobel 105 and North 1996). Models of photosynthetic carbon gain also predict a low proportion of root biomass and shallow rooting for systems with pulsed water availability, across plant 106 phenotypes (Schwinning and Ehleringer 2001). However, much of this knowledge is 107 108 based on studies of desert succulents. Sedum species used on green roofs are usually from 109 less extreme environments, where one would expect more flexible strategies for resource 110 acquisition, as reflected in their facultative photosynthetic C₃-CAM metabolism (Winter and Holtum 2014). Although spatial patterns of soil nutrients trigger morphological root 111 112 foraging responses in many species (Kembel and Cahill 2005), such responses have, to 113 the best of our knowledge, not been investigated in Sedum species. More knowledge on 114 this part could give input to how to place fertilisers to direct rooting patterns on green roofs. Interestingly, strong root foraging for Cd and Zn have been found for Zn/Cd 115 116 hyperaccumulating genotypes of Sedum alfredii (Liu et al. 2010).

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To obtain relevant knowledge for use in green roof systems, we addressed some of these questions in an experiment under greenhouse conditions. The objective of the study was to evaluate the extent to which *Sedum* species are able to exploit water and nutrients by 121 root foraging in substrates of different composition and layering. We tested the hypotheses that i) Sedum species actively forage for soil resources, resulting in a higher 122 123 root density in substrate layers with more nutrients or higher water retention capacity, and 124 ii) Root placement is determined by the water availability of the substrate layers, so foraging in layers with high water-holding capacity is weakened when water availability 125 is increased through watering. As a consequence, more root biomass and root length 126 127 would be allocated to deeper layers in a layered substrate when fertiliser or water-holding 128 material is placed at the bottom. In sum, these tests can also inform whether substrate 129 modifications that can improve shoot survival would have negative impact on the root 130 foraging for resources.

131

132 Materials and methods

The interactive effects of substrate texture, layering, irrigation, and fertiliser placement
on root foraging were tested for the species *Sedum album* (L.) and *S. rupestre* (L.) in a
greenhouse pot experiment during June-September 2016.

136

137 Substrate texture

138 We used four substrate compositions: a fine substrate, a coarse substrate, a mixed

substrate as a 1:1 combination of fine and coarse material, and a layered substrate with

140 the coarse mixture on top of the fine mixture (Fig. S1). All four substrates were based on

141 different fractions of pumice that were initially sieved to fine (0-2 mm), intermediate (2-

142 5 mm) and coarse (5-10 mm) fractions and then combined to a fine (40% fine + 34% intermediate fractions) and a coarse (26% intermediate + 48% coarse fractions) base 143 144 mixture. These base mixtures were combined with 9% sieved mature and nutrient-poor compost and 17% gravel (3-5 mm). All proportions are by volume, and all final substrates 145 were blended for 2 minutes in a concrete mixer. We used 11 cm tall square pots (10 cm 146 by 10 cm) filled to 9 cm with substrate. This corresponds well with the recommended 147 148 thicknesses of extensive green roof substrates and these small pots were used to simulate 149 the rapid fluctuations in water content on green roofs. Total pore volume was 42 and 46 % 150 and maximum water capacity 0.5 and 0.33 kg water per L substrate for the fine and coarse components, respectively. Substrate pH measured in a 1:5 solution with distilled water 151 ranged from 7.5 to 7.6. 152

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154 Watering and fertiliser placement

Fertiliser placement and watering regime were varied while keeping the other of the two 155 156 factors constant. For the watering regime comparison, all pots had fertiliser evenly mixed 157 throughout the substrate depth. All pots received 1.0 g of granular Multicote 4 slowrelease fertiliser (15-7-15 + Micronutrients, Haifa Chemicals Ltd.), designed to release 158 159 nutrients over a 4 month period at 21 °C. The pots received three irrigation regimes, with 160 weekly individual watering from the top applying 50, 100 or 200 mL per pot using tap water of low conductivity (0.15 mS cm⁻¹), corresponding to 5, 10 or 20 mm water depth 161 162 per week. For the fertiliser placement experiment, the fertiliser was mixed into the substrate either in the top or bottom halves of the pots, or evenly into the substrate of the 163 164 whole pot. Fertiliser placement was only manipulated for pots receiving the 100 mL week⁻ ¹ watering regime. Pots were placed in random positions on a net frame on a greenhouse
table for unrestricted drainage.

167 Vertical water distribution was documented in pots without plants, by weighing and drying samples of substrate of the middle upper and middle lower parts of pots for each 168 169 substrate combination. Pots were tilted to remove water laying on the inside of the pot 170 and samples taken 10 minutes after water addition. Water content of the substrates was 171 0.2 g/g before testing and samples were dried at 105 °C for 24 h before weighing. Pots retained almost all the water at 5 mm. Pots were saturated at about 5 (coarse), 10 172 173 (layered and mixed) and 20 mm (fine) for the respective substrates. (Fig. S2). The fine 174 substrate consistently retained more water in the upper part than the other substrate, in 175 addition to retaining more in total. The mixed and layered pots retained about the same amount of water with a similar partitioning, except the layered substrate retained more 176 177 in the bottom half for the 5 mm treatment. In coarse substrates, 50 % or more of 178 retained water was retained in the lower half of the pot.

179

Plants received only natural irradiance and during the experiment they experienced mean diurnal temperature of 18.7 °C (95% confidence interval 18.4-19.2 °C). Mean diurnal minimum air temperature of 15.8 and maximum of 25.1°C gave a night drop of 10.3°C on average. Temperature extremes were maximum 34.8 and minimum 12.2°C. Over the experimental period, the plants experienced an approximate 1031 growing degree-days over a base temperature of 10 °C.

186 Reference evapotranspiration (ET_0) was estimated using the Penman-Monteith equation

187 (FAO-56) and summed over weekly intervals according to the watering schedule.

188 Estimated cumulative weekly ET_0 was well above 5 mm, except for the last week of the

experiment, and above 10 mm for the first 7 weeks (Fig. S3). It was never above 20 mm
per week. The study site in SW Norway is characterised by a cool, wet maritime climate
(Köppen-Geiger, Cfb). During the past 20 years, the summer period (May-August) has
had 19% of weeks with less than 5 mm, 29% with less than 10 mm and 47% with less
than 20 mm of accumulated precipitation. Hence, the given watering treatments
correspond well with the drier parts of the growing season in the region, also
representative of the original locations of the plant material.

196

197 Plant material

Small plug plants of *Sedum album* (L.) and *S. rupestre* (L.) propagated from cuttings were 198 199 used in the experiment originating from populations in Southern Norway. S. album is part of the Leucosedum clade within the Crassulaceae (van Ham and 't Hart 1998), while S. 200 201 rupestre belongs to the Rupestria series, often raised to the rank of a separate genus, 202 *Petrosedum* (Mort et al. 2001) and more closely related to *Sempervivum* than to *S. album*. 203 Thus, these two species span some of the variation within the polyphyletic 'Sedum' genus. 204 The plants were established in 4 cm deep pyramidal plugs of a coarse material similar to 205 the substrates used in the experiment for 8 weeks until the experiment and the plugs were 206 rooted. Shoots of transplants were 30-40 mm long and had a biomass of 42 ± 8 (SD) mg for S. album and 71 ± 10 mg for S. rupestre. Root fractions of the total biomass were 0.1 207

- and 0.2 respectively. To ease transplantation, the experimental pots were watered daily
- 209 for a week after planting before the experimental treatments started.

211 Harvests

At harvest after 12 weeks, shoots were cut at the surface, dried for 48 h at 70 °C and 212 213 weighed. The pot substrate was cut in half at the interface of the coarse and fine mixtures 214 or at the same depth for the other substrates. Roots were washed out of each pot half, 215 scanned using a calibrated dual-light flatbed scanner (Epson Perfection V700 Photo 216 Scanner, Epson America Inc., CA, USA) and analysed for total root length and root 217 diameter using the WinRhizo software (Regent Instruments Inc., Québec, Canada). After 218 scanning, root biomass was dried and weighed as for shoots. Care was taken to analyse 219 roots only and not buried parts of stems. 220 A foraging index was calculated for each pot as the difference in root length (FIRL) or 221 root biomass (FIBM) between the upper and the lower half of the pot, divided by the total 222 root length or total root biomass per pot. A high value of FIRL or FIBM (i.e. values close

to 1.0 (or -1.0)) indicates a strong bias towards root development in the upper (or lower) half of the pot, while a value close to zero indicates that root development is similar throughout the substrate depth. The root fraction of the total biomass (R_f) was calculated as the ratio of root biomass to total biomass per pot.

To check root distribution within pots, the soil from frozen pots with *S. album* was cutin three horizontal layers, and each layer cut in 16 even sized cubes. Roots were washed from these cubes and root biomass determined. This was done for the mixed substrate and 10 mm watering only (Fig. S4).

231

232 Experimental design and statistical analyses

233 We used a design with two species by four substrate structures by three watering regimes or fertiliser placements by four replicates, giving 96 pots per experiment and 160 pots in 234 total, with 32 pots common to both experiments. The effects of watering and fertiliser 235 236 placement were analysed separately in 3-way ANOVA models using the general linear model option in Minitab 17 (Minitab Ltd., Coventry, UK), with species, substrate 237 structure and the water or fertiliser treatments as fixed factors. Model diagnostics were 238 239 evaluated using QQ plots of residuals and plots of residuals against predicted values. Two outliers for root length and root biomass were identified by their strongly deviating length 240 241 to biomass ratios and were replaced with treatment means. Partial effect sizes were estimated as ω_P^2 (Olejnik and Algina 2003). ANOVA results and effect sizes were used 242 to identify important results, where only significant effects with a considerable effect 243 sizes were considered major effects. 244

245

246 **Results**

247 Overall growth patterns

Starting with about the same transplant biomass, the species had average relative growth rates over 12 weeks of between 0.046 and 0.060 g g⁻¹ day⁻¹ for *S. album* and 0.032 and 0.052 g g⁻¹ day⁻¹ for *S. rupestre*. The corresponding mean increase in total biomass was between 2.0 and 7.0 g per plant and between 1.1 and 5.5 g per plant, respectively. Both species had an allocation of biomass to roots of 5-13% of total biomass (Fig. 1). Specific root length varied between 200 and 265 m g⁻¹ and root length per shoot biomass varied between 10 and 24 m g⁻¹. Both estimates were affected by substrate structure, but did not

- differ between species (Table 1, Fig. 2). Growth was vegetative during the whole
- 256 experiment.

Table 1. Effects of watering regime or vertical fertiliser placement on growth responses of two *Sedum* species (*S. album* and *S. rupestre*) to

substrate structure and layering. F and P values from ANOVA models are shown with effect sizes, estimated as partial ω^2 . Error df = 72,

total df = 95. Major effects evaluated by the P values and the effect sizes are indicated in bold.

		Total root length			Shoot biomass			Root biomass (LN)			Root fraction (R _f)			Specific root length		
Source	df	F	Р	ω_{P}^{2}	F	Р	ωP ²	F	Р	ω_P^2	F	Р	ω_P^2	F	Р	ω_P^2
Effects of watering																
Species	1	64.46	0.000	0.40	75.33	0.000	0.44	32.34	0.000	0.25	3.47	0.066	0.03	2.56	0.114	0.02
Structure	3	8.08	0.000	0.18	31.02	0.000	0.49	6.95	0.000	0.16	20.29	0.000	0.38	11.22	0.000	0.25
Water	2	45.27	0.000	0.48	187.43	0.000	0.79	59.65	0.000	0.55	2.79	0.068	0.04	13.70	0.000	0.21
Sp*Str	3	4.07	0.010	0.09	3.43	0.021	0.07	3.13	0.031	0.06	1.43	0.241	0.01	0.53	0.660	-0.02
Sp*W	2	0.05	0.950	-0.02	0.28	0.754	-0.01	1.46	0.238	0.01	0.54	0.586	-0.01	1.06	0.353	0.00
St*W	6	1.93	0.088	0.06	4.60	0.001	0.19	0.82	0.556	-0.01	1.26	0.287	0.02	1.53	0.180	0.03
Sp*St*W	6	1.09	0.376	0.01	0.88	0.511	-0.01	1.16	0.336	0.01	1.50	0.190	0.03	1.15	0.340	0.01
\mathbb{R}^2 adj			66			86			65			42			39	
Effects of fe	ertiliser	placement														
Spec	1	49.24	0.000	0.34	190.40	0.000	0.67	55.89	0.000	0.37	3.96	0.050	0.03	0.00	0.981	-0.01
Structure	3	0.62	0.605	-0.01	61.60	0.000	0.66	4.83	0.004	0.11	16.00	0.000	0.32	2.14	0.102	0.03
Fertiliser	2	0.52	0.597	-0.01	1.95	0.149	0.02	0.24	0.787	-0.02	0.14	0.867	-0.02	0.13	0.882	-0.02
Sp*St	3	1.72	0.170	0.02	0.58	0.629	-0.01	3.23	0.027	0.07	3.72	0.015	0.08	1.27	0.290	0.01
Sp*F	2	0.37	0.693	-0.01	3.65	0.031	0.05	1.40	0.253	0.01	0.49	0.614	-0.01	0.85	0.434	0.00
St*F	6	1.39	0.229	0.02	8.72	0.000	0.33	2.68	0.021	0.10	4.11	0.001	0.16	0.99	0.437	0.00
Sp*St*F	6	1.75	0.121	0.05	2.69	0.021	0.10	1.52	0.184	0.03	0.83	0.547	-0.01	0.61	0.723	-0.03
R ² adj			36			82			47			43			0	

		Foraging	g index roo (FIRL)	t length	Foraging index root biomass (FIBM)			Root length per shoot biomass			Root diameter bottom			Root diameter top		
	df	F	Р	$\omega_{\rm P}^2$	F	Р	$\omega_{\rm P}^2$	F	Р	$\omega_{\rm P}^2$	F	Р	$\omega_{\rm P}^2$	F	Р	$\omega_{\rm P}^{2}$
Effects of w	atering															
Species	1	30.52	0.000	0.24	29.10	0.000	0.23	0.12	0.732	-0.01	3.84	0.054	0.03	0.27	0.607	-0.01
Structure	3	2.40	0.075	0.04	1.65	0.186	0.02	21.42	0.000	0.39	4.22	0.008	0.09	1.42	0.243	0.01
Water	2	12.32	0.000	0.19	20.81	0.000	0.29	21.18	0.000	0.30	3.59	0.033	0.05	0.15	0.861	-0.02
Sp*St	3	0.59	0.621	-0.01	0.34	0.796	-0.02	1.41	0.246	0.01	0.23	0.873	-0.02	0.15	0.929	-0.03
Sp*W	2	1.03	0.361	0.00	0.31	0.736	-0.01	1.13	0.328	0.00	3.59	0.033	0.05	2.90	0.061	0.04
St*W	6	1.74	0.123	0.04	3.23	0.007	0.12	1.86	0.099	0.05	1.91	0.090	0.05	0.86	0.526	-0.01
Sp*St*W	6	0.56	0.758	-0.03	0.39	0.883	-0.04	0.33	0.920	-0.04	0.80	0.571	-0.01	1.71	0.131	0.04
R ² adj.			38			44			52			21			4	
Effects of fe	rtiliser	placement														
Spec	1	0.86	0.357	0.00	17.19	0.000	0.15	1.02	0.316	0.00	2.95	0.090	0.02	1.33	0.253	0.00
Structure	3	3.39	0.022	0.07	8.12	0.000	0.18	12.00	0.000	0.26	5.48	0.002	0.12	2.00	0.121	0.03
Fertiliser	2	4.69	0.012	0.07	35.82	0.000	0.42	0.55	0.579	-0.01	2.05	0.136	0.02	0.44	0.643	-0.01
Sp*St	3	0.51	0.674	-0.02	1.73	0.169	0.02	0.31	0.820	-0.02	0.77	0.516	-0.01	1.64	0.188	0.02
Sp*F	2	1.05	0.354	0.00	1.30	0.279	0.01	0.47	0.628	-0.01	2.63	0.079	0.03	1.64	0.202	0.01
St*F	6	0.38	0.888	-0.04	0.78	0.586	-0.01	1.16	0.338	0.01	2.41	0.036	0.08	2.57	0.026	0.09
Sp*St*F	6	0.38	0.887	-0.04	1.33	0.254	0.02	0.65	0.693	-0.02	0.23	0.964	-0.05	1.35	0.245	0.02
R^2 adj.			6			54			23			1	6		16	

269 Effects of watering regime

Both substrate structure and watering had large effects on plant growth, while interactions
between them were few (Table 1). Shoot and root biomass and total root length increased
with watering (Fig. 3), while the root fraction of the total biomass was not affected. Both
the specific root length and root length per shoot biomass decreased with watering (Fig.
and the root length per shoot biomass was considerably lower in the fine substrate (Fig.
2).

276 Although the interactive effect of watering and substrate structure and layering on shoot biomass was significant (Table 1), the responses to watering followed similar patterns in 277 278 all substrates, only with a slightly stronger response to watering in the fine (S. album) and 279 fine and mixed (S. rupestre) substrates (Fig. 3). The two species had different growth 280 responses to substrate structure, but these differences were not affected by watering (species by structure vs. species by structure by water interactions, Table 1). Sedum album 281 was less able to exploit the deeper layers of the layered substrate, expressing similar shoot 282 283 biomass and root length as for the coarse mix (Fig. 3).

284

285 Effects of fertiliser placement

Overall, fertiliser placement had weaker effects on plant growth than watering and no effects on shoot and root biomass, total root length and root fraction were found (Table 1, Fig. 3). The effect of fertiliser placement on shoot biomass differed between substrates (Table 1), primarily as a consequence of a more positive effect of fertiliser placement near the top of the substrate in the fine substrate. There were no major differences in shoot 291 biomass in response to fertiliser placement between species (despite the significant species by fertiliser interaction, Table 1). Top fertilisation also gave higher root biomass 292 293 in the fine and mixed substrates; while an even fertiliser distribution gave more root 294 biomass in the layered substrate. The interaction between structure and fertiliser 295 placement for the root fraction (Table 1) was due to higher R_f for even fertiliser 296 distribution in the layered substrate and lower R_f for even fertiliser distribution in the 297 mixed structure (not shown). In summary, combining fertiliser and the fine substrate in 298 the bottom layer did not increase root biomass or root length there compared to the other 299 configurations.

300

301 Effect of substrate structure

Layered, mixed and fine substrates all gave higher shoot and root biomass than the coarse 302 303 substrate, and the fine substrate gave higher shoot biomass than layered and mixed substrates (Table 1, Fig. 3). Combined, this meant that plants growing in mixed, layered 304 and coarse substrates had a higher proportion of their total biomass (R_f) allocated to roots 305 306 than plants in the fine substrate (Fig. 2). Plants in the fine substrate also had considerably lower root length per shoot biomass. The specific root length was higher in the coarse and 307 mixed than in the layered and fine substrates, accompanied by slightly thinner roots in 308 309 the coarse and mixed substrates (Table 1, 0.36-0.37 mm compared with 0.38-0.39 mm). 310 Substrate structure had no effect on root diameter in the upper half of the pot, but the 311 layered substrate gave thicker roots in the bottom half of the pot than the coarse and mixed 312 substrates for both the water and fertiliser experiments (Table 1). The layered substrate gave a root diameter increase in the lower part of the pots, from 0.33-0.34 to 0.37 mm for 313

S. album and from 0.34 -0.35 to 0.38 mm for *S. rupestre*, but these differences are small
as also reflected in the small effect sizes (Table 1).

316

317 Root foraging

Increased watering reduced the allocation of root length and root biomass to the lower part of the pots (Fig. 5, Table 1). This effect differed between substrates, with a more negative effect of watering on biomass allocation to the lower part of pots in the layered and mixed substrates (Fig. 5). In contrast, the response in root length allocation to watering was not affected by the substrate composition (Table 1).

Although fertiliser placement had a significant effect on the foraging index of root length (FIRL), this effect was marginal (Table 1, Fig. 5). This corresponds with the weak responses of root length to fertiliser placement and substrates. Root biomass, however, followed the placement of the fertiliser to a larger extent than root length (Fig. 5, Table 1). Placement of fertiliser in the bottom half of pots increased the allocation of root biomass in this part (and lowered the FIBM). This effect was not dependent on substrate structure (Table 1).

330

Both foraging indexes showed a positive relationship with shoot biomass in the water dataset for both species, while there were no such relationships in the fertilizer dataset (not shown). Breaking down these relationships on treatments and species, there were no consistent patterns.

335

336 **Discussion**

337 Our main hypothesis was that *Sedum* roots show active foraging for water and nutrients. As we found significant responses in root foraging to both watering and fertilisation 338 treatments, this hypothesis was not rejected. However, although we found some flexibility 339 340 in root allocation patterns, the ability for precise root foraging was low to moderate, as also reflected in the low proportion of biomass allocated to roots. Hence, these Sedum 341 342 species had only a limited ability to exploit resources like water and nutrients by precise 343 root foraging in substrates of different composition and layering within the 3-months 344 timeframe of this experiment. Water was the factor driving plant growth, followed by 345 substrate structure, while fertiliser placement had only a marginal effect on plant growth.

346

347 Overall effects of substrate structure

348 Across treatments, substrate structure affected many components of plant growth. The 349 main distinction was between the fine substrate and the others, where fine substrate gave a higher shoot biomass, a lower root fraction and more shoot biomass per unit root length. 350 351 This finding is in line with the better water-holding capacity of the fine substrate, 352 providing water for a longer period between the weekly watering (Fig. S2). The coarse substrate also differed from the layered and mixed substrates for some responses, in 353 354 principle reflecting the same mechanisms, but at the other end of the humidity gradient. 355 Except for the 5 mm watering, the coarse pots retained about half the amount of water as 356 the mixed and layered pots (Fig. S2). Positive relationships between water-holding 357 capacity of the substrate and plant performance have been documented in several studies. It has been shown that thicker substrates (Durhman et al. 2007; Ondoño et al. 2016), 358 substrates with finer particles (Raimondo et al. 2015), substrates with water-holding 359

additives (Savi et al. 2014) and substrates with more organic matter (Nagase and Dunnett
2011) improve plant growth and/or survival across different environmental conditions.
The results for the fine substrate fit well with these findings.

The layered substrate improved plant performance compared with the coarse substrate. 363 364 Based on standardised tests, the coarse substrate was able to hold 330 g of water, the layered substrate 420 g and the fine substrate 500 g per litre of substrate. The realised 365 water retention was considerably lower with about 50, 100 and 200 g per pot of about 0.5 366 L (Fig. S2), the difference caused by different compaction and the time allowed for water 367 368 absorption. Considering the strong response to watering and the differences in biomass 369 between the layered and the fine substrate, it is noteworthy that this substantial increase 370 in available water in the layered compared with the coarse substrate was not fully exploited. 371

372 With the low proportion of root biomass, Sedum contributions to carbon sequestration 373 will primarily be through aboveground biomass. Our estimates of the biomass fraction in 374 roots is lower than found by Getter et al (2009), but clearly there are large differences 375 between succulent species where the deciduous *Phedimus* species had a higher potential 376 for C binding in roots (Getter et al 2009). Long-term effects need to take root turnover and degradation into account. Considerably better alternatives than *Sedum* based roofs 377 378 exists for carbon sequestration, like more diverse green roofs and ground based solutions (Whittinghill et al. 2014). 379

380 Effects on root foraging

381 The effect of substrate structure differed between watering and fertiliser placement382 treatments and affected primarily shoot biomass and the root fraction of the total biomass.

383 However, we found no interactions between substrate structure and fertiliser placement on the foraging indexes FIRL or FIBM and only a weak interaction between substrate 384 385 structure and watering level for FIBM. Although the hypothesis of that the effect of substrate composition on root foraging would depend on fertiliser placement and/or 386 watering level could not be rejected, there was no solid support for it. Accordingly, we 387 388 found no strong support for the prediction that more root biomass and root length would 389 be allocated to deeper layers in a layered substrate when fertiliser or fine material is placed 390 at bottom. Fertiliser placement in the bottom half of the pots increased root biomass there, 391 but this effect was independent of substrate structure. Fine material both holds more water and has the potential to retain more nutrients than the coarse material. Interactions 392 393 between water and nutrients have been found in other systems where root biomass follows 394 both water and nutrient placement (Wang et al. 2007). We used a nutrient-poor compost 395 to add some organic material to the substrates. Although this was leached for soluble 396 nutrients before use, it released some nutrients to the plants throughout the experiment and masked some of the effects of fertiliser placement. In conclusion, nutrient availability 397 did not limit plant growth, so a strong root foraging for nutrients could not be expected. 398 399 Coarse green roof substrates leach considerable amounts of nutrients (Kuoppamäki and 400 Lehvävirta 2016), but that would depend on the precipitation or as in our case the watering treatments. This interaction between watering and fertiliser placement was not included 401 402 in the experimental design.

403

Fig. 3 indicated more shallow roots in the layered substrate (higher FIRL) at increasing
watering. This effect could be interpreted as a weakening of foraging in layers with higher
water-holding capacity when water availability is increased through watering. However,

this was a common trend for most of the substrates (also with a main effect of watering)
showing just a more shallow rooting at increasing watering. As we found no preferential
foraging in specific layers, we were not able to evaluate the hypothesis that increased
watering reduced the foraging in substrate layers with higher water-holding capacity

411

Except for the fine substrate, there were only weak effects on the root fraction of the total biomass. The overall patterns of root and shoot growth and allocation of biomass to roots in response to watering reflected those found for *Sedum lineare* under different watering regimes (Lu et al. 2014). This indicates that there is a limited flexibility in the allocation of biomass to roots, even under water-deficient conditions also in other *Sedum* species.

The lack of interactions between most treatments on root foraging is difficult to explain, 417 418 especially the observation that roots did not forage deeper in layered substrates at the lowest watering level, where weekly watering was below ET₀ throughout most of the 419 experimental period. Growth was clearly water-limited, as shoot biomass increased by 51 420 and 152 % when going from 5 to 10 and 20 mm week⁻¹, respectively. There are some 421 422 alternative explanations. Either the soil water conditions were not extreme enough to 423 trigger a change in rooting patterns, or morphological root plasticity in response to 424 especially water availability is not a common strategy in *Sedum* species. Rooting depth is a plastic trait in many plants, and non-succulent species respond to early signals of soil 425 426 drying (Schachtman and Goodger 2008) by allocating resources to deeper roots (Comas 427 et al. 2013). We do not know if root elongation in *Sedum* species is more or less sensitive 428 to soil water potential than that in non-succulents. Observations that succulent species 429 can extend their roots in dry soil with water from the shoot (North and Nobel 1998) 430 indicate that they may be less sensitive. Recent findings have shown the importance of shoot-derived abscisic acid (ABA) for root growth (McAdam et al. 2016). As the 431 432 succulent leaves of *Sedum* species are buffered against loss of turgor for extended periods 433 during drought (Sayed et al. 1994), one can speculate on the extent of signalling from shoots to roots before leaf turgor decreases. CAM species can be considered to show 434 hypersensitivity to ABA and rapidly respond to environmental conditions to conserve 435 436 water (Negin and Moshelion 2016). This indicates that strategies to prevent losses are 437 more important than foraging.

438

439 Succulents are somewhat difficult to classify using the competitor-stress tolerator-ruderal 440 (CSR) model of primary plant strategies developed by Grime and colleagues (Hodgson et al. 1999), but Sedum species are considered stress-tolerators. There are trade-offs 441 442 among strategies, so stress-tolerant species in less productive systems and in systems 443 where abiotic constraints dominate are less likely to express costly foraging strategies based on changes in morphology, relying instead on cellular acclimations (Grime and 444 Mackey 2002). Such trade-offs lower the root foraging precision and competitive ability. 445 446 Sedum species have been found to perform well even on substrates as thin as 2.5 cm 447 (Durhman et al. 2007), although without competition they perform better on thicker 448 substrates (Getter KL, Rowe 2008; Thuring et al. 2010) and substrates with higher water retention capacity (MacIvor et al. 2013). However, there are some species-specific 449 450 responses and differences between broadleaved (like *Phedimus*) and 'cylindrical' Sedum species (MacIvor et al. 2013). 451

452

453 Justification of the approach

Duration of the experiments is one critical factor when evaluating allocation strategies. 454 455 During the 3-months experiment, plants experienced 1031 growing degree days (with a base temperature of 10 °C), showed a 73 to 107 fold increase in shoot biomass in S. album 456 457 and 23 to 45 fold increase in S. ruprestre, and had a total root length at harvest ranging 458 of from 6.4 to 8.2 m in S. album and from 3.8 to 5.9 m in S. ruprestre. In our opinion, 459 there was sufficient time and growth to detect flexibility in rooting patterns. These patterns may however change over time and there may be seasonal patterns in root growth 460 461 strategies not detected in our study. These aspects have not been documented for *Sedum* 462 species so far and critical factors as root turnover and expected lifetime of Sedum roots 463 are unknown. As nutrients did not limit plant growth, the test for root foraging for nutrients is weak and should be followed up by more studies. 464

Pot size is another critical factor, causing edge effects and restricts access to resources. 465 At start, the pots had a total plant biomass to rooting volume ration (BVR) of from 0.05 466 to 0.08 gL⁻¹. At harvest, this had increased to an average of 3.4 gL⁻¹ (95% CI of 3.1 to 3.8, 467 range 0.7 to 7.4). This is higher than 1 gL^{-1} as recommended for pot experiments by 468 Poorter et al (2012), but considerably lower than for established green roofs. Using data 469 470 from Getter et al. (2009), considering only aboveground biomass and a mean carbon content of 42 %, twelve standard Sedum based green roofs had a mean BVR of 8.7 gL⁻¹ 471 $(95\% \text{ CI: } 6.2 - 11.2 \text{ gL}^{-1})$. In small pot volumes, root foraging along pot walls is common 472 473 and roots are usually forced downwards when they meet the pot wall. This would however obscure the rationale of our approach. Previous observations of Sedum root development 474 475 in these media do however predicted a more homogenous root distribution.

e observed a rather homogenous horizontal root distribution, and not a higher root density
along pot edges (Fig. S4). This is as expected with such porous substrates and illustrates
that the edge effects were small. In conclusion, the chosen pot size was suitable to
represent the extensive green roof systems studied with respect to both available soil
volume and the rapid changes in soil water content on green roofs.

481 Conclusions

482 Both *Sedum album* and *S. rupestre* showed a low ability to exploit water and nutrients by precise root foraging in substrates of different texture and layering. Allocation of biomass 483 to roots was low and showed limited flexibility, even under water-deficient conditions. 484 More shallow roots were produced at higher irrigation and in fine substrate. However, 485 486 considerably more shoot biomass developed per unit root length in fine substrate. A 487 layered substrate with coarse substrate on top of a layer of fine substrate did not give major improvements compared with a coarse or a mixed substrate, and led to no additional 488 foraging of root biomass or root length in the deep layer, even when fertiliser was placed 489 490 in this layer. Thus layered substrates provide no major additional benefits for Sedum 491 growth and roof function during summer. This also infers that it will be difficult to direct roots to deeper layers, at least in the short term. A stronger response to fertilizer placement 492 493 is however expected when nutrients are more limiting. In summary, water was the main 494 factor driving plant growth, followed by substrate structure, while vertical fertiliser 495 placement had marginal effects on plant growth.

496

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Fig. 1 Biomass allocation patterns in *Sedum album* and *S. rupestre* in response to watering (5, 10 or 20 mm week⁻¹) and fertiliser placement (top half, bottom half or evenly distributed in pots) when cultivated in green roof substrates of contrasting structure. Note that root data are multiplied by a factor of 10 for better presentation



Fig. 2 Specific root length, root length per shoot biomass and root fraction (mean with 95% confidence interval) for *Sedum album* (white symbols) and *S. rupestre* (black symbols) growing in green roof substrates of contrasting composition. Estimates are averaged over watering and fertilisation treatments



Fig. 3 Effects of watering, vertical fertiliser placement and substrate structure on shoot biomass, total root length and total root biomass (mean \pm SD) in *Sedum album* and *S. rupestre* growing in green roof substrates of contrasting texture



Fig. 4 Specific root length and root length per shoot biomass (mean with 95% confidence interval) of *Sedum* species growing in green roof substrates of contrasting composition receiving 5, 10 or 20 mm water per week. Estimates are averaged over species (*S. album* and *S. rupestre*)



Fig. 5 Effects of watering (above) and vertical fertiliser placement (below) on indices of root foraging (mean with 95% confidence interval) based on root length (FIRL) or root biomass (FIBM) for two *Sedum* species grown in substrates of contrasting texture, receiving either 5, 10 or 20 mm irrigation per week or manipulation of vertical fertiliser placement in the pots (B = bottom, M = mixed, T = top). Indices were estimated as response in upper part of pot minus response in bottom part of pot divided by the sum

response for the whole pot. The dashed lines indicate when root length or root biomass is evenly distributed between the top and bottom parts of the pot

SUPPLEMENTARY MATERIAL

Sedum root foraging in layered green roof substrates

Peng Ji¹, Arne Sæbø², Virginia Stovin³, Hans Martin Hanslin^{2*}

* Corresponding author: hans.martin.hanslin@nibio.no



Fig. S1 Texture and layering in the four types of substrates used in the pot experiment. The full height of the columns corresponds to the substrate height of 9 cm used in the pots. Black bar is 1 cm.



Fig. S2 The amounts of water lost and retained in the different vertical layers given 5, 10 or 20 mm of watering. The bottom figure shows the relationship between water added and water retained per pot (mean \pm SD, n=3). Stippled line is the 1:1 relationship between added and retained.

Amount watering (g)



Fig. S3 Estimated cumulative weekly reference evapotranspiration (ET_0) during the experiment. Dashed lines show the irrigation regimes of 5 and 10 mm week⁻¹



Fig. S4 Contour plots showing the horizontal distribution of root biomass for three layers of the pot volume in a mixed substrate estimated as mean percentage (%) of root biomass per horizontal layer based on a sampling of 16 cubes per layer (n = 3). Pot base is 9 by 9 cm.