

1 Release of Bet v 1 from birch pollen from 5 European 2 countries. Results from the HIALINE study

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59 **Abstract**

60

61 Exposure to allergens is pivotal in determining sensitization and allergic symptoms in
62 individuals. Pollen grain counts in ambient air have traditionally been assessed to
63 estimate airborne allergen exposure. However, the exact allergen content of ambient
64 air is unknown. We therefore monitored atmospheric concentrations of birch pollen
65 grain and the matched major birch pollen allergen Bet v 1 simultaneously across
66 Europe within the EU-funded project HIALINE (Health Impacts of Airborne Allergen
67 Information Network).

68 Pollen count was assessed with Hirst type pollen traps at 10 l/min at sites in France,
69 United Kingdom, Germany, Italy and Finland. Allergen concentrations in ambient air
70 were sampled at 800l/min with a Chemvol high-volume cascade impactor equipped
71 with stages $PM > 10\mu m$, $10\mu m > PM > 2.5\mu m$, and in Germany also $2.5\mu m > PM > 0.12\mu m$. The major birch pollen allergen Bet v 1 was determined with an
73 allergen specific ELISA. Bet v 1 isoform patterns were analyzed by 2D-SDS-PAGE
74 blots and mass spectrometric identification. Basophil activation was tested in an
75 Fc ϵ R1-humanized rat basophil cell line passively sensitized with serum of a birch
76 pollen symptomatic patient.

77 Compared to 10 previous years, 2009 was a representative birch pollen season for
78 all stations. About 90% of the allergen was found in the $PM > 10\mu m$ fraction at all
79 stations. Bet v 1 isoforms pattern did not varied substantially neither during ripening
80 of pollen nor between different geographical locations. The average European
81 allergen release from birch pollen was 3.2 pg Bet v 1/pollen and did not vary much
82 between the European countries. However, in all countries a >10-fold difference in
83 daily allergen release per pollen was measured which could be explained by long-
84 range transport of pollen with a deviating allergen release. Basophil activation by
85 ambient air extracts correlated better with airborne allergen than with pollen
86 concentration.

87 Although Bet v 1 is a mixture of different isoforms, its fingerprint is constant across
88 Europe. Bet v 1 was also exclusively linked to pollen. Pollen from different days
89 varied >10-fold in allergen release. Thus exposure to allergen is inaccurately
90 monitored by only monitoring birch pollen grains. Indeed, a humanized basophil
91 activation test correlated much better with allergen concentrations in ambient air than
92 with pollen count. Monitoring the allergens themselves together with pollen in
93 ambient air might be an improvement in allergen exposure assessment.

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98 Introduction

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100 Allergies are the most prevalent chronic disease in Europe with an >20% prevalence
101 (Bauchau and Durham, 2004; Sunyer et al., 2004; Bousquet et al., 2007). Allergic
102 diseases to airborne allergens have been steadily increasing over the past decades
103 (Eder et al., 2006). This increase in prevalence is also due to replacement of older
104 less sensitized individuals in the population by younger individuals with a higher
105 degree of sensitization (Jarvis et al., 2005; Rönmark et al., 2009; Laatikainen et al.,
106 2011). For developed countries a leveling-off of allergic disease prevalence was
107 reported (Asher et al., 2006; Grize et al., 2006), but for some highly developed
108 countries like Finland an end of this epidemic is not in sight (Laatikainen et al., 2011).
109 Allergen exposure determines sensitization (Olmedo et al., 2011) and allergy
110 symptoms correlated with exposure (Corrigan et al., 2005; Brito et al., 2011).
111 Exposure to outdoor airborne allergens is monitored by determining the concentration
112 of pollen in ambient air with a network of over 350 pollen traps spread over Europe
113 (www.polleninfo.org, accessed January 2012). However, humans react to the
114 allergen and the concentration of airborne pollen (the pollen count) is a proxy of
115 exposure. Indeed, several investigations imply that the pollen count might not be
116 representative for allergen exposure, also because allergen was found in non-pollen
117 bearing fractions of ambient air (Schäppi et al., 1997b; De Linares et al., 2010;
118 Fernandez-Gonzalez et al., 2011).

119 Pollen grains release more immunologically active compounds than only allergen,
120 like PALMS, adenosine and NADPH oxidase (Dharajiya et al., 2007; Gilles et al.,
121 2009; Gilles et al., 2011). These compounds can act as adjuvants, however the
122 allergen from pollen is the dominant factor for evoking symptoms (Brito et al., 2011).

123 Almost all patients allergic to birch pollen are allergic to Bet v 1, sometimes
124 accompanied by a sensitization to Bet v 2 or Bet v 4 (Moverare et al., 2005). The Bet
125 v 1 content of birch pollen is not constant (Buters et al., 2010), and geographical
126 variation was described (Buters et al., 2008). Climate change with increasing
127 concentrations of CO₂ results in higher pollen production as CO₂ is both an airborne
128 fertilizer and a greenhouse gas. Changes in allergen release per pollen would be in
129 addition to the changed load of airborne pollen (Estrella et al., 2006; Rogers et al.,
130 2006; Shea et al., 2008; Ziska and Beggs, 2012). We therefore determined the
131 variation in the release of the major birch pollen allergen Bet v 1 with an
132 immunochemical ELISA method in the project HIALINE (Health Impacts of Airborne
133 Allergen Information Network) and confirmed this independently in selected cases
134 with a bio-assay using FcεR1-humanized rat basophils. We evaluated whether the
135 used methods, Chemvol® and ELISA, were suited for an allergen-release measuring
136 network. We also investigated whether meteorological factors could govern allergen
137 release from pollen, in an effort to predict the effect of climate change on the
138 allergenicity of pollen.

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Materials and methods

Pollen count

Airborne concentrations of pollen were sampled with volumetric spore traps of the Hirst design (Hirst, 1952) and examined by light microscopy as described before (Smith et al., 2009). Pollen counting methods vary historically between countries and three different methods were included in this study. Three stations used a technique where slides were examined along three (France) or four (Germany and Italy) longitudinal transects (Sikoparija et al., 2011). In the UK pollen grains were counted along twelve latitudinal transects (Smith et al., 2009). The analysis method used in Finland is random sampling of microscopic fields which has been shown to give parallel results to the counts carried out with the two other methods (longitudinal and latitudinal transects) (Mäkinen, 1981). Difference between methods was eliminated by correction for surface counted (Comtois et al., 1999). Pollen counts for the 10-years average were obtained from the EAN (European Aeroallergen Network, <https://ean.polleninfo.eu>, accessed January 2012). The pollen season was calculated as described in the legend of Table 1. The pollen index, the total exposure to pollen from one season at one station was expressed as the sum of daily average pollen concentrations according

$$\int_{t=0}^{t \rightarrow \text{end}} C_t \cdot d(t)$$

and carries the unit Σ pollen grains/m³.

Quality control of the pollen count was monitored by the UK partner. All partners filled in questionnaires requesting data on the siting and operation of the pollen trap and the preparation and counting of samples. In addition, calibration slides were sent to each station and everyone involved in counting pollen for the study examined the slides. The limits imposed on the quality control survey were: (1) pollen counts between 0-30 pollen grains/m³ had to be within ± 10 pollen grains/m³; (2) pollen counts >30 pollen grains/m³ had to be within $\pm 30\%$. The rule that pollen count between 0-30 pollen grains/m³ had to be within ± 10 pollen grains/m³ was introduced because very low pollen count can easily vary by more than 30% (Sikoparija et al., 2011). This number ($\pm 30\%$) was also determined by Comtois et al. as the inherent variation of the method (Comtois et al., 1999).

The variability between Hirst type volumetric spore traps was determined with 3 samplers operating simultaneously over a 3-week period at <5m apart on a rooftop at 9m a.s.l. during the birch pollen season in Munich, Germany (n=63, pollen between 0 and 4500 grains/m³).

Airborne allergen sampling

Air was sampled as previously described (Buters et al., 2010). In brief: 800l/min ambient air was sampled on polyurethane foam with a high-volume Chemvol® cascade impactor equipped with size class stages PM>10 μ m and 10 μ m>PM>2.5 μ m

185 (Butraco Inc., Son, Netherlands) (Demokritou et al., 2002). In Munich, the stage
186 $2.5\mu\text{m} > \text{PM} > 0.12\mu\text{m}$ was also sampled. Air flow was kept constant with a rotameter
187 controlled high-volume pump (Digitel DHM-60, Ludesch, Austria). At each site, the
188 Chemvol® sampler was located at equal height and within 5m of a Hirst-type trap.
189 For each station Chemvol® and Hirst type pollen samples were analyzed daily for
190 identical time periods. Polyurethane foam impacting substrates were cut into 3
191 identical parts per day and stored at $\leq -20^\circ\text{C}$ until extraction.

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194 *Meteorological data*

195 Weather data (daily average temperature, daily average humidity, cumulative daily
196 rainfall, average wind speed, and cumulative daily sunshine) were measured using
197 sensors at the stations. Missing parameters were obtained from the closest nearby
198 stations of the national weather services. In the UK this was the Pershore station, in
199 France the station at the airport of Bron, in Germany station 3379, Munich City of the
200 Deutsche Wetterdienst, in Finland it was Turku Artukainen (Airport) weather station.
201 In Italy all was measured at the same location as the Chemvol sampler.

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204 *Extraction and analysis of Bet v 1*

205 The Chemvol® polyurethane impacting substrates were extracted for 4h in 0.1M
206 ammonium bicarbonate pH8.1 in a head-over-head rotator. Extracts were aliquoted,
207 frozen, lyophilized and stored at $\leq -20^\circ\text{C}$. Aliquots were reconstituted in 0.1M
208 phosphate buffered saline pH7.4, serial diluted and allergen was determined by using
209 a 2-site binding assay based on monoclonal antibodies in an ELISA format.
210 Monoclonal antibodies 4B10 and 2E10 specific for Bet v 1, natural purified Bet v 1
211 standards and controls were provided by our partner Allergopharma KG, Reinbek,
212 Germany (Chapman et al., 2008; Kahlert et al., 2008).

213 With each ELISA two control samples of different concentration were analyzed. The
214 values of these controls had to be within 25% of a reference value for the ELISA to
215 be accepted. Then only those values of serial dilution that yielded the same
216 concentration were reported.

217 For each day at least two filter parts were independently analyzed. If two filter parts
218 did not yield a value within 25% of each other, a third extraction was performed and
219 analyzed. The final reported concentration of each day was the mean of all valid
220 determinations, mostly the mean of at least 16 ELISA wells. The same Standard
221 Operating Procedure (SOP) was used by all partners, which included written data
222 inclusion rules.

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225 *Bet v 1- isoform analysis*

226 Pollen was sampled sequentially from several locations across Europe (not always
227 the stations where the allergen measurements were done) as described previously by
228 Buters et al. (Buters et al., 2010) and shipped at -20°C to our partner at
229 Allergopharma. Pollen from the day of pollination of each *Betula pendula* tree were
230 extracted (see above) and subjected to 2D-SDS-PAGE electrophoresis using a first
231 dimension of isoelectric focusing between pH4-7 (IPG strips, GE Healthcare, Munich)
232 and a second dimension of SDS-PAGE (Excel SDS gel, 12 – 14%, GE Healthcare,
233 Munich) for size separation. Gels were stained with Coomassie Brilliant Blue, blotted

234 and dried. Available recombinant Bet v 1.0401 (Bet v 1d) and Bet v 1.0601 (Bet v 1f)
235 were a kind gift of Prof. F. Ferreira, University of Salzburg, Austria, recombinant Bet v
236 1.0101 was from Allergopharma. Spots were quantified using Proteomweaver
237 software (Definiens, Munich, Germany) and expressed as relative % of the sum of all
238 intensities. Punched spots were identified by tryptic digestion and analysis by mass
239 spectrometry as described by Sarioglu et al. (Sarioglu et al., 2008).

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242 *Degranulation of humanized RBL*

243 The humanized rat basophil cell line RBL-2H3 clone h21 expressing the α , β , and γ
244 chain of the human Fc ϵ R1 was a kind gift of Prof. S. Vieths, Paul Ehrlich Institute,
245 Langen, Germany. Cells were grown under standard conditions (Vogel et al., 2005)
246 and were passively sensitized with serum of a birch pollen symptomatic patient (skin
247 prick test positive and RAST>3). Dilutions of daily Chemvol samples were added to
248 the cells and degranulation was quantitated as β -hexosaminidase release,
249 determined as nitrophenol release from pNAG (p-nitrophenol-D-2-acetamido-2-
250 deoxyglucopyranosid, Sigma-Aldrich Corp, St. Louis, MO) in relation to total β -
251 hexosaminidase activity after lysis of the cells with 1% Triton-X100 (Vogel et al.,
252 2005). Only values within the linear dose-response range of the cells (5-45%
253 degranulation) were reported. Because extracts vary greatly in Bet v 1 content,
254 degranulation was calculated as if 1m³ air was given to the cells. This could result in
255 hypothetical degranulations of up to 800%.

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258 *Modeling*

259 The System for Integrated modeLLing of Atmospheric coMposition (SILAM, (Siljamo
260 et al., 2008a)) was used to compute the footprints of the observations and also to
261 evaluate the flowering season.

262 A footprint of a single observation is, by definition, a surface area that delineates the
263 sources that are responsible for the observed atmospheric concentrations.
264 Interpreted in probabilistic terms, the footprint shows the probability of a specific air
265 parcel to reach the observational site. The areas, for which this probability is not zero,
266 comprise the footprint of the particular observation. As a simplification, all sources
267 located within the footprint area would influence this observation, and no sources
268 located outside the footprint area would affect it. The specific contributions of these
269 sources vary depending on the footprint value: the higher the value, the stronger the
270 source impact. Computations of the footprint with standard dispersion models are
271 prohibitively resource-consuming, while the adjoint modeling used in this study
272 generates the solution with reasonable efforts (Sofiev et al., 2006b).

273 The flowering season prediction followed the thermal-type model as described before
274 (Sofiev et al., 2006a; Siljamo et al., 2008b). The SILAM model was run with a time
275 step of 15 minutes and evaluated the transport for 60 hours backward in time, for
276 each daily observation at each site. The configuration included 8 vertical layers up to
277 ~6 km above the ground. The horizontal grid cell size was 25 km and the domain of
278 simulations covered almost the whole of Europe. Meteorological information was
279 taken from the operational archives of the European Centre of Medium Range
280 Weather Forecast (ECMWF). This data had a spatial resolution of about 25km and
281 time step of 3 hours.

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Statistical analysis

Differences were analyzed with a paired Student's *t*-test unless stated otherwise (Livingston, 2004). A *p* value <0.05 was considered statistically significant. Outliers were defined as more than 3 standard deviations of the mean. The relationship between allergen and pollen count was presented graphically by scatter plot. The strength of the relation was expressed by the coefficient of correlation (r^2) which was calculated by using linear regression (Lorenz, 1989). The same pre-given spreadsheet was used for all calculations.

293 Results

294

295 *Pollen count*

296 Although the pollen counting method differed between the stations, this was
297 compensated by correcting for the counted surface area and reporting the number as
298 pollen grains/m³. This is a common method also used by EAN. Indeed, when we
299 counted the same slides for birch pollen both latitudinal or longitudinal (n=15) in the
300 same laboratory by the same operator this resulted in a <7% difference, in
301 agreement with the literature for other pollen species (Carinanos et al., 2000).

302 In our network out of a total of 28 calibration counts for *Betula*, three were outside the
303 limits imposed on the Quality Control survey (11%).

304 The variability of pollen counts at the same location between 3 Hirst type pollen traps
305 was 23%. Recounting pollen from the same slide (n=8) by the same operator showed
306 a <4% variability in counting reproducibility, the same as reported before (Kapyla and
307 Penttinen, 1981). Thus 19% of the variation in birch pollen count is due to differences
308 between the samplers.

309

310 The annual sum of pollen (pollen index, see methods) in 2009 varied between 235 Σ
311 pollen grains/m³ in Italy to 3144 Σ grains/m³ in Germany (see Table 1). The average
312 annual birch pollen count in 2009 was 55% of the 1999-2009 average birch pollen
313 flights for these stations (range 30 to 90%, see Table 1), and similar years did occur
314 for each station (data not shown). Thus the birch pollen season in 2009 was
315 representative for all stations. Other European stations (not in this manuscript) report
316 higher 10-year average counts for birch pollen (i.e. central Finland, Poland or
317 Ukraine), as the center of birch tree habitat is the eastern part of Europe just outside
318 the European Union (www.polleninfo.org, accessed January 2012). Thus our results
319 cover the extremes of the European Union habitat for birch trees.

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321

322 *Airborne pollen allergen*

323 The detection limit of the ELISA was 0.7 ng Bet v 1/ml, which equals 2.1 pollen/m³.
324 Running a high and a low control concomitant with each run monitored the
325 performance of the Bet v 1-ELISA. Data inclusion rules were installed. The variability
326 of the ELISA (n=66, all stations, see Table 2) of the low control (1.9 ng/ml) was
327 17.4% and 12.8% for the high control (7.4 ng/ml), in close agreement with the
328 literature (Schäppi et al., 1996; Buters et al., 2010). The pollen count and allergen
329 Bet v 1 concentrations are depicted in Figure 1. Allergen was 89.6±1.5% found in the
330 PM>10μm fraction, the remainder was in the 10μm>PM>2.5μm fraction, and none in
331 the smallest 2.5μm>PM>0.12μm fraction, available only in Munich. No allergen was
332 found when no pollen was detected (see Figure 1). The European average Bet v 1
333 release per pollen was 3.2 pg Bet v 1/pollen (r²=0.714, see Figure 2). However, the
334 difference in allergen release per pollen between days and locations was >10-fold,
335 even if we deleted all pollen count <10 pollen/m³ to avoid high allergen release
336 values due to less reliable (low) pollen count. When we removed (arbitrarily) the Bet v
337 1 release values per pollen stemming from pollen count <10 pollen/m³, then the
338 average allergen release per pollen of the 10% lowest values was 0.61 pg Bet v
339 1/pollen, the average of the highest 10% released 8.76 pg Bet v 1/pollen. Within each

340 station, the average allergen release of the highest 5% and the lowest 5% values
341 also varied at least >10-fold (see Figure 2). Between countries the average allergen
342 release per pollen were considered similar as the observed differences are within the
343 uncertainties in pollen count (method variation <30%) or ELISA determination
344 (method variation <17%).

345

346

347 *Bet v 1 isoforms*

348 Bet v 1 is increasingly expressed during ripening in the last week before pollination
349 (Buters et al., 2010). The pattern of Bet v 1 isoforms during ripening was determined
350 using one tree, (see Figure 3). Spots of April 18th were identified using mass
351 spectrometry and, if available, by using pure recombinant isoforms.

352 The isoform pattern of Bet v 1 did not differ markedly during ripening, (see Figure 3).
353 Thus the same fingerprint of isoforms is expressed at rising concentrations during
354 pollen ripening. An exception is spot 1, which was analyzed as being a truncated
355 isoform of Bet v 1.0101 (Bet v 1a), which increases upon ripening of the pollen.
356 However, this was a minor Bet v 1 isoform.

357 The ELISA antibody combination recognized all isoforms equally as pooled human
358 serum. The ELISA antibodies did not recognize the isoform Bet v 1.0401 (Bet v 1d),
359 which was also not recognized by pooled human serum from 10 donors (data not
360 shown) (van Ree et al., 2008). Our ELISA thus represents human reactivity.

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362 The isoform pattern of birch pollen from several different locations across Europe
363 varied marginally, see Figure 4. Because ripening of the pollen does not influence the
364 isoform pattern, harvesting at not exactly the same time points before pollination
365 would not explain differences. We conclude that the Bet v 1 isoform pattern is the
366 same during ripening and the same across Europe. Our antibodies recognized all
367 isoforms except the hypoallergenic isoform Bet v 1.0401, like humans (data not
368 shown) (see Figure 3) and the difference in Bet v 1 content between the stations is
369 thus due to differences in amount of released Bet v 1, not due to release of different
370 isoforms.

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373 *Degranulation of basophils*

374 Rat basophils carrying the human FcεR1-receptor were passively sensitized with
375 serum of a birch pollen sensitive individual and incubated with daily extracts of
376 Chemvol samples from Munich, Germany. The reactivity of the cells is depicted in
377 Figure 5. Again, degranulation was only seen when pollen were counted. In addition,
378 β-hexosaminidase release correlated well ($r^2=0.95$, insert in Figure 5a) with Bet v 1
379 concentration in ambient air, but less well with pollen count from Munich ($r^2= 0.71$,
380 see insert in Figure 5b). The level of detection, defined as 10% degranulation above
381 baseline, was 0.2 ng/ml Bet v 1.

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383

384 *Modeling*

385 For this analysis, due to uncertainties in pollen counts and allergen determination, we
386 omitted pollen counts below 15pollen/m³ and allergen content below 10pg/m³. Above
387 this values the measurements are more robust. Analysis of the time series of the
388 allergen release per pollen grain shows several features: (i) an established value for

389 allergen release per pollen, which is steadily between 2 and 4pg/grain during the
390 whole season for all stations, (ii) small fluctuations around this value, which are
391 probably due to uncertainty of both Bet v 1 and pollen observations, (iii) several
392 episodes at some stations lasting for a few days, during which the allergen content
393 differs from the average level by several-fold.

394 By joint analysis of the observation footprints and flowering patterns, the allergen
395 content during the multi-day episodes can be correlated with geographical location of
396 the pollen sources, as shown in Figure 6. The allergen content observed in Turku is
397 systematically low during the period 12 - 17 May, with the gradual decrease during
398 12-15 May down to less than 1 pg/pollen and then increase again during 15-17 May.
399 The footprint analysis showed that the transport direction of pollen was gradually
400 changing from central Sweden to northern Finland and then further to southern
401 Finland. All these areas were flowering during these days. This suggests that the
402 pollen originating from northern Finland showed an about three-times lower allergen
403 release than that from the more southern regions. A similar pattern was seen again
404 24 and 25 May when the footprint was covering northern Sweden, which was
405 flowering at that time (not shown). There was again a 3 times lower allergen release
406 in pollen from the north than during the days when pollen originated from more
407 southern regions.

408 In Munich, the episode during April 16-18 was characterized with high allergen
409 content, flanked before and afterwards by several days with an average allergen
410 release per pollen. Before and after that period the footprint shows the source areas
411 to the east and somewhat to the south of Munich and the allergen release was at the
412 average level. However, during the episode, the pollen mainly originated from
413 mountains of Switzerland, where birch was at full flowering. Those grains had 2-3
414 times more allergen release than regions to the east of Munich. The part of footprint
415 covering the Alps was probably void because there was no flowering in the high
416 mountains yet.

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420 Discussion

421
422 Birch pollen and the major birch pollen allergen Bet v 1 were sampled during the
423 birch pollen season in France, United Kingdom, Germany and Finland (and some
424 data from Italy). The birch pollen season in 2009 was a representative year for all
425 stations.

426 We did not find any large difference in Bet v 1 isoform patterns during ripening of the
427 pollen or across Europe, except for a minor isoform (truncated Bet v 1.0101, less
428 than 5% of total Bet v 1). The isoform pattern recognized by ELISA was identical to
429 the pattern recognized by a serum pool of birch allergic individuals. The average Bet
430 v 1 release per pollen from the different stations was within the error of the methods
431 for pollen counting (30%) and ELISA (17%) and we consider the Bet v 1 release from
432 pollen in 2009 similar across Europe. However, daily Bet v 1 release varied >10-fold
433 in all countries. This was independently confirmed with a bio-assay for Bet v 1 using
434 human sIgE in humanized rat basophils. Indeed, β -hexosaminidase release in these
435 cells as a proxy for histamine release correlated better with Bet v 1 than with pollen
436 concentrations.

437 We conclude that birch pollen can vary >10-fold in allergen release and that this
438 variation is similar across Europe.

439

440

441 *Pollen count*

442 Although pollen counts in Europe have been performed using standard methods for
443 decades (Hirst, 1952), little is known about the reproducibility of Hirst type volumetric
444 spore traps (Barral et al., 2009). In Munich, birch pollen count recorded by three
445 volumetric pollen traps simultaneously varied by 19% for birch pollen. Comparing the
446 different pollen counting methods (longitudinal, latitudinal and random) by sending
447 calibration slides to all stations showed that the majority of birch pollen count (89%)
448 varied by less than 30%, in agreement with previous publications (Comtois et al.,
449 1999; Carinanos et al., 2000; Sikoparija et al., 2011).

450 Comtois et al. suggested that there will always be imprecision linked with the airborne
451 pollen count unless aerobiologists count the whole slide, and even then variation
452 between pollen traps will still be present. There is always a trade off between
453 precision and the amount of time required to produce the daily pollen count (Comtois
454 et al., 1999).

455 Based on these results and the literature we believe that a variation of <30% in our
456 data for pollen count is already accounted for by the pollen sampling and counting
457 methodology and is not due to variations in nature.

458 The pollen season in 2009 was a representative year for all stations, as the average
459 annual sum of birch pollen in 2009 was 55% (range 30-90%) of the 10-year average
460 for each station with no outliers (see Table 1). The stations also cover a larger
461 geographic area within Europe.

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463

464 *Allergen Bet v 1 in ambient air*

465 The pH of the extraction buffer is likely to influence the amount and isoform pattern of
466 the allergens extracted. We used a slight alkaline extraction buffer, a condition

467 observed in nasal fluid of allergic patients (Podoshin et al., 1991) and recommended
468 by Cadot et al. for the appearance of relevant isoforms (Cadot et al., 1995). Indeed,
469 extraction at pH8.1 was used by several, but not all authors (Rantio-Lehtimäki et al.,
470 1994; Schäppi et al., 1997b; Petersen et al., 2001).

471
472 The average Bet v 1 release across Europe was 3.2 pg Bet v 1/pollen and is similar
473 to the value of 4 pg Bet v 1/pollen reported before (Schäppi et al., 1997b). Schenk et
474 al. also published that the variation in allergen release between birch species was
475 limited (Schenk et al., 2011) but report higher allergen content per pollen, which
476 could be due to their extraction method. Another difference with other authors is that
477 in the current study pollen were sampled from ambient air and not from trees or
478 commercial suppliers.

479 The average Bet v 1 release per pollen per country did not vary much (-23.1% in the
480 UK and +21.8% in Germany, see Figure 2) and could be explained by variations in
481 methodology as we determined that between pollen samplers a variation of 23%
482 existed and our ELISA has a <17% variation between laboratories in agreement with
483 the literature where 30% variations for pollen monitoring is reported (Comtois et al.,
484 1999; Sikoparija et al., 2011) and similar variations were found for ELISA allergen
485 determinations (Schäppi et al., 1996; van Ree et al., 2008).

486 However, when comparing the 10% lowest allergen release values per pollen with
487 the 10% highest values, the daily allergen release difference across Europe and also
488 within each country was >10-fold. Several other authors also report differences in
489 allergen release per pollen for birch and olive pollen (Pehkonen and Rantio-Lehtimäki,
490 1994; Schäppi et al., 1997a; De Linares et al., 2007; Buters et al., 2010; Brito et al.,
491 2011). This study shows that the variation in allergen release per pollen is substantial
492 but equally distributed across the European birch tree habitat.

493 Across Europe, about 90% of the allergen was recovered from the >10 μ m fraction,
494 none in the 2.5 μ m>PM>0.12 μ m fraction (only in Munich), and no allergen was
495 detected when no pollen was detected, neither with our ELISA nor with a more
496 sensitive bio-assay. This shows that the only source of allergen is birch pollen, in
497 agreement with previous results where allergens were monitored for several years on
498 a row at one location (Buters et al., 2010). Birch allergen containing particles were
499 reported in the fractions PM<10 μ m (birch pollen have a geometric diameter of 21-
500 24 μ m (Brown and Irving, 1973; Rantio-Lehtimäki et al., 1994; Schäppi et al., 1999))
501 indicating in combination with our results that if such particles exist, their appearance
502 is rare and might need specific atmospheric conditions like thunderstorms (D'Amato
503 et al., 2008), which did not occur during our experiments.

504

505

506 *The role of Bet v 1 isoforms*

507 We found that the Bet v 1 isoform fingerprint was more or less constant during
508 ripening and across Europe with minor variations (see Figures 3 and 4), in agreement
509 with other authors (Friedl-Hajek et al., 1999; Eler et al., 2011; Schenk et al., 2011).
510 The isoform truncated Bet v 1.0101 did increase in expression upon ripening of the
511 pollen, however this minor truncated Bet v 1.0101 isoform represented <5% of total
512 Bet v 1 isoforms. The identity of the spots was confirmed with LC-MS after tryptic
513 digest (Sarioglu et al., 2008), and if possible, by comparison with recombinant
514 proteins. Also, our antibodies did not recognize the hypoallergenic isoform Bet v
515 1.0401 (Bet v 1 d), similar to a pool of human serum (data not shown) and as

516 reported by others (Friedl-Hajek et al., 1999; Erler et al., 2011). Thus the difference
517 we measured must be due to different total concentrations of allergenic Bet v 1 of
518 which 50-70% is Bet v 1.0101(Bet v 1a) (Erler et al., 2011).

519
520

521 *Clinical relevance*

522 We used an independent method from ELISA to determine allergen content in
523 ambient air by taking human serum from a birch pollen sensitive individual and
524 passively sensitizing basophils (see Figure 5). Although human mast cells would be
525 the preferred cells, few human mast cell lines are available and none has been
526 shown to be suitable for allergen determination (Kirshenbaum et al., 2003; Guhl et al.,
527 2010). Besides mast cells, basophils are also responsible for the clinical symptoms of
528 allergy in humans (Gibbs, 2007). The basophil cell line with human Bet v 1 specific
529 IgE as detector of environmental Bet v 1, resulted in an immune response as seen
530 with allergic individuals (Vogel et al., 2005). Although more tedious, this method is
531 more sensitive than the ELISA, able to detect 0.2 ng Bet v 1/ml (defined as
532 degranulation >10%). With this more sensitive bio-assay also no allergen was
533 detected when no pollen were measured. Also, β -hexosaminidase release (a
534 substitute marker for histamine release which is the hallmark of allergic disease),
535 correlated well with Bet v 1 ambient concentrations ($r^2=0.95$), but lesser good with
536 pollen count ($r^2=0.72$, see Figure 5).

537
538

539 *Modeling*

540 We found several multi-day episodes where the differences in pollen potency could
541 be explained by differences in origin of the pollen (see Figures 1 and 6). We omitted
542 pollen counts below 15 pollen/m³ and allergen measurements below 10 pg/m³ to
543 guarantee more robust data. We also focused on multi-day episodes, as one-day
544 jumps of the pollen content are more difficult to analyze. In general, there can be
545 three possible explanations for single day jumps:

546 (i) similar to multi-day episodes, the peaks may correspond to specific
547 transport conditions and/or origin of the grains. However, footprint analysis
548 did not reveal such dependence.

549 (ii) low pollen count means higher uncertainty of the allergen release per
550 pollen. However, it should manifest itself as both anomalously high and low
551 values, which is not the case: almost all low-count cases were
552 characterized by the high allergen release.

553 (iii) there can be allergen present in air apart from the one encapsulated in the
554 pollen grains (D'Amato et al., 2008). Even when the number of grains is
555 low, this extra allergen could lead to high release estimates. The
556 instrumentation used in the current study does not detect such allergen,
557 thus does not allow an explicit check of this possibility. Using suitable
558 equipment, no such free allergen was detected (Buters et al., 2010).

559

560 Several multi-day episodes were detected (see Figure 6). They corroborate our
561 finding with ELISA and the bio-assay that pollen is not constant in their allergen
562 release.

563

564 Noteworthy, the average value also suggests some north-to-south gradient: for
565 Munich and Worcester the allergen release is about 3 pg/pollen, whereas for Turku it
566 is about 2 pg/pollen (statistically non significant).

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568

569 *Conclusions*

570 In daily samples taken during the birch pollen season in 5 European countries we
571 could only detect allergen when pollen was present. Also allergen was predominantly
572 found in the PM_{>10}µm fraction, the fraction where the pollen land in the cascade
573 impactor. Thus Bet v 1 in ambient air was confined to birch pollen. With our method
574 the average European allergen release per pollen was 3.2pg Bet v 1/pollen. The
575 average allergen release in 2009 did not vary substantially between countries.
576 However, a >10-fold difference between daily allergen release per pollen was
577 detected in all countries. Thus pollen exposure qualitatively represents allergen
578 exposure but not quantitatively. The allergen concentration also correlated better with
579 the bio-assay for immune response than pollen concentration. Modeling showed that
580 multi-day episodes exist were pollen from specific origins consistently varied in
581 allergen release. Thus we expect allergen monitoring to be a more accurate predictor
582 of human allergic symptoms than pollen count.

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584

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602 Legend to the Figures

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604 **Figure 1.** Daily values in 2009 for birch pollen (grey bars) and Bet v 1 (colored lines)
605 in PM_{>10}μm (green) and 10μm>PM_{>2.5}μm (red) from the different European
606 stations. (A) Munich, Germany (B) Turku, Finland (C) Lyon, France (D) Worcester,
607 UK. Only Munich, Germany additionally sampled 2.5μm>PM_{>0.12}μm (yellow). Note:
608 the scales differ between stations for clarity. The amount of daily allergen released
609 per pollen is given for pollen counts >10 pollen/m³ (pink).

610

611

612 **Figure 2.** Correlation between the total releasable allergen in the air and pollen count
613 in the different countries in 2009. The slope of each linear regression curve
614 represents the average Bet v 1 release per pollen for that country. Each point
615 represents duplicate determinations of one day in one country. - ● - Germany; - ■ -
616 Finland; - ▲ - France; - ▼ - United Kingdom; - ◆ - Italy. The data point indicated with
617 the arrow was treated as an outlier.

618

619

620 **Figure 3.** One D (A) and 2D (B) SDS-PAGE blots of 0.1 M ammonium bicarbonate
621 extracts from pollen harvested sequentially in 2008 from one tree in Germany. April
622 19-20 were days of maximum pollination of that tree. Lane A is the image of the blot
623 of the molecular weight marker (15kDa) and pollen extract after separating according
624 to size before development into the second pH4-7 dimension in the right lanes.
625 Identity of the spots was determined for April 18th. Spot 0101 represents Bet v
626 1.0101(Bet v 1a), the others are Bet v 1.0401(Bet v 1d), Bet v 1.0601 (Bet v 1f), Bet v
627 1.1401 (Bet v 1m) and truncated Bet v 1.0101(t0101, see methods).

628

629

630 **Figure 4.** Isoforms of Bet v 1 from trees across Europe in 2009. The same
631 methodology as for Figure 3 is used.

632

633

634 **Figure 5.** β-hexosaminidase release (a proxy of histamine release) of FcεR1-
635 humanized RBL cells after passive sensitization with serum of a birch pollen
636 sensitized individual (see methods) and exposure to daily samples of TUM2009.
637 Samples were diluted to fit the dynamic range of the cells. Degranulation per cubic
638 meter air was then calculated and depicted and may exceed 100%. (A) Bet v 1
639 concentrations and β-hex- (B) Pollen concentration and β-hexosaminidase release.

640

641

642 **Figure 6.** Flowering of birch trees (pink) and the observation footprint (area where
643 the particles collected in the instrument stem from considering the last 60hrs, blue).
644 Date, pollen concentration and their potency (pg Bet v 1/pollen) is given. The potency
645 of pollen can depend on the area of origin. (A) Turku, Finland and (B) Munich,
646 Germany.

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