ComparISON OF BUFFERs FOR EXTRACTION OF 
MITE ALLERGEN DER P 1 FROM DUST

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Der p 1 is the main allergen of house dust mite Dermatophagoides pteronyssinus, which has routinely been detected in residential dust. However, the procedure for extracting Der p 1 from reservoir dust has not been well defined. The aim of this study was to compare Der p 1 mass fractions in dust extracts prepared using the following extraction buffers: phosphate (pH 7.4), borate (pH 8.0), and ammonium bicarbonate (pH 8.0), all with 0.05 % Tween 20. Twenty-eight dust samples were divided into three aliquots and each portion was extracted with one of the three buffers at room temperature. Der p 1 mass fractions were measured in a total of 84 dust extracts using the enzyme immunoassay (range: 0.1 μg g⁻¹ to 7.53 μg g⁻¹). Statistical methods including intraclass correlation showed a high agreement between Der p 1 mass fractions irrespective of the extracting medium. Our results suggest that all three buffers are suitable for the extraction of mite allergens and routine Der p 1 analysis in dust.

KEY WORDS: Dermatophagoides pteronyssinus, ELISA, extraction buffers, indoor allergens, intraclass correlation, settled dust

House dust mites are the source of 21 allergens identified so far. The major allergens of the common dust mite (Dermatophagoides pteronyssinus) are Der p 1 and Der p 2 (1-3). Exposure to these allergens is associated with allergic symptoms and asthma in sensitised people (1-3). Reservoir dust samples have been used as a proxy for Der p 1 exposure in residential (1, 4) and occupational (5) settings. In order to compare results from different studies investigating allergen exposure and related health effects, both the collection techniques and laboratory protocols (analysis, extraction, and storage) should be comparable. Enzyme-linked immunosorbent assay (ELISA) is the standard method for quantification of common indoor allergens in reservoir dust (6). However, there is no standard protocol for the extraction of allergens from dust (7). Several solutions have regularly been used for extraction of Der p 1 in laboratories worldwide including phosphate, borate, and ammonium bicarbonate buffers (Table 1). Some laboratories add the non-ionic surfactant Tween 20 to the extraction medium and some do not. Little information is available about the extraction efficiency of buffers on Der p 1 measurements. Siebers et al. (8) found that the type of buffer affected measurement of Der p 1 levels. In this short report, Der p 1 concentrations (high exposure level) in a borate extract were much higher than in phosphate and ammonium bicarbonate extracts. However, other operating conditions (such as time and temperature) appeared to be very important for Der p 1 extraction and Der p 1 measurement with ELISA. Extraction at a lower temperature (4 °C) resulted in lower Der p 1 level, irrespective of the buffer type. This lack of standardised operating conditions, certainly contributes to great inter-laboratory differences in Der p 1 measurements (7, 9).
The aim of our study was to establish correlations between Der p 1 mass fractions in dust samples extracted with three common buffers, namely phosphate, borate, and ammonium bicarbonate, all containing 0.05% Tween 20, at room temperature.

MATERIALS AND METHODS

Dust collection

Twenty-eight dust samples were collected from 18 urban households in Zagreb, Croatia between 2007 and 2009. Samples were taken by vacuuming a carpeted area of the living rooms using a standard vacuum cleaner adapter and cellulose filter (Heska AG, Freiburg, Switzerland) as described earlier (10).

Dust extraction and analysis

Three buffers were used for Der p 1 extraction from settled dust: phosphate (PBS; pH 7.4), borate (BBS; pH 8.0; Titrisol, Merck, Germany), and 0.125 mol L⁻¹ ammonium bicarbonate (ABS; pH 8.0; Kemika, Zagreb). The final concentrations of the PBS components were 137 mmol L⁻¹ for NaCl, 10 mmol L⁻¹ for Na₂HPO₄·2H₂O, 2 mmol L⁻¹ for KH₂PO₄, and 2.7 mmol L⁻¹ for KCl. According to the manufacturer (Merck, Germany), the borate buffer consisted of 0.11 mol L⁻¹ H₃BO₃, 0.044 mol L⁻¹ HCl, and 0.056 mol L⁻¹ NaOH. All buffers contained 0.05%

<table>
<thead>
<tr>
<th>Extraction buffer</th>
<th>Tween 20</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate (PBS) pH 7.4</td>
<td>/</td>
<td>23, 24</td>
</tr>
<tr>
<td>PBS²</td>
<td>0.05</td>
<td>20, 25</td>
</tr>
<tr>
<td>PBS (1 h, 30 °C)</td>
<td>0.05</td>
<td>26</td>
</tr>
<tr>
<td>PBS (2 h, RT)</td>
<td>0.05</td>
<td>27, 28, this work</td>
</tr>
<tr>
<td>PBS-1 % BSA (2 h, RT)</td>
<td>0.05</td>
<td>29</td>
</tr>
<tr>
<td>PBS-1 % BSA (overnight, RT)</td>
<td>0.5</td>
<td>30</td>
</tr>
<tr>
<td>PBS-0.2 % BSA (overnight, 4 °C)</td>
<td>0.2</td>
<td>31</td>
</tr>
<tr>
<td>Borate (BBS) pH 8.0</td>
<td>/</td>
<td>13, 32-35</td>
</tr>
<tr>
<td>BBS³</td>
<td>0.05</td>
<td>this work</td>
</tr>
<tr>
<td>BBS-5 % BSA³</td>
<td>/</td>
<td>36</td>
</tr>
<tr>
<td>BBS-5 % BSA (overnight, 4 °C)</td>
<td>/</td>
<td>37</td>
</tr>
<tr>
<td>BBS-aprotinin (2 h, 4 °C)</td>
<td>0.1</td>
<td>38</td>
</tr>
<tr>
<td>Ammonium bicarbonate (ABS) pH 8.0</td>
<td>/</td>
<td>39</td>
</tr>
<tr>
<td>ABS (2 h, RT)</td>
<td>/</td>
<td>21, this work</td>
</tr>
</tbody>
</table>

RT - room temperature
BSA - bovine serum albumin
⁻ - extraction conditions not available

<table>
<thead>
<tr>
<th>Der p 1</th>
<th>Median</th>
<th>Mean±SD</th>
<th>Range</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>low level</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS-T</td>
<td>0.648</td>
<td>0.721±0.53</td>
<td>0.12 to 1.655</td>
<td>18</td>
</tr>
<tr>
<td>BBS-T</td>
<td>0.753</td>
<td>0.696±0.497</td>
<td>0.105 to 1.62</td>
<td>18</td>
</tr>
<tr>
<td>ABS-T</td>
<td>0.645</td>
<td>0.616±0.462</td>
<td>0.060 to 1.565</td>
<td>18</td>
</tr>
<tr>
<td>moderate level</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PBS-T</td>
<td>3.620</td>
<td>3.774±1.681</td>
<td>2.015 to 7.255</td>
<td>10</td>
</tr>
<tr>
<td>BBS-T</td>
<td>3.605</td>
<td>3.958±1.685</td>
<td>2.075 to 7.42</td>
<td>10</td>
</tr>
<tr>
<td>ABS-T</td>
<td>3.422</td>
<td>3.838±1.667</td>
<td>2.270 to 7.525</td>
<td>10</td>
</tr>
</tbody>
</table>

PBS-T - phosphate buffer-Tween
BBS-T - borate buffer-Tween
ABS-T - ammonium bicarbonate buffer-Tween
Tween 20 (T) (Merck, Germany). Before extraction, each sample was manually sieved through a 300 μm sieve, mixed until homogenous, and weighed. Fine dust samples were divided into three 100-mg aliquots, and 2 mL of extraction solution was added to each aliquot. Extractions were done at room temperature with constant shaking on a Vortex mixer (Ika Vortex, Germany) for 2 h. After 10 min of centrifugation at 1,000x$g$, supernatants were stored in plastic tubes at -20 °C until analysis for Der p 1 content. A total of 84 dust extracts were analysed for Der p 1 content.

The mass fractions of Der p 1 were determined with capture ELISA, using a commercial kit (Indoor Biotechnologies Ltd, Cardiff, UK) as described in our earlier article (10). The kit contained monoclonal antibody SH8 (mouse anti-Derp 1, IgG2A) (lot number 30034) as capture antibody, biotinylated monoclonal antibody 4C1 (mouse IgG1; lot number 30068) as secondary antibody, and Der p 1 standard (2500 ng mL⁻¹). All antibodies (except capture mAb), standards, dust extracts, and positive and negative controls were diluted in PBS-T containing 1 % bovine serum albumin (BSA; PBS-T-BSA; Sigma, USA). Dust extracts were diluted three or six times depending on the Der p 1 level. Aliquots of extracts were placed into a 96-well microtitre plate (Maxi Sorp, Nunc, Denemark) following the manufacturer’s instructions. After all reagent incubations, optical densities were read at 450 nm using a microtitre plate ELISA reader (IASON, Vienna, Austria). The limit of detection was 0.1 μg g⁻¹. The intra-assay coefficient of variation (CV) and inter-assay CV for Der p 1 ELISA were 6.9 % and 13.1 %, respectively (10).

### Statistical analysis

All data were analysed using free statistical software R, version 2.13.2. Descriptive statistics was used to illustrate the distribution of Der p 1 in PBS-T, BBS-T, and ABS-T extracts. According to allergen mass fraction, Der p 1 values were grouped in low (from 0.1 μg g⁻¹ to 2.0 μg g⁻¹) and moderate (from 2.01 μg g⁻¹ to 7.53 μg g⁻¹) levels. Der p 1 measurements were compared for each pair of buffers and, in a separate procedure, for all three buffers. For each pair of buffers, Der p 1 measurements were compared using correlation coefficients (Pearson’s and Spearman’s), and linear regression coefficients (slope and intercept, with 95 % confidence intervals, 95 % CI) in two mass ranges. The agreement between Der p 1 values for each pair of buffers was estimated using the intraclass correlation coefficient (ICC) (11, 12). This coefficient was also used to determine the agreement between Der p 1 values in all three buffers, since other coefficients allow only paired measurements.

Intraclass correlation coefficients were calculated from two-way random effects model with both buffers and dust batches as covariates (11) using R software. Intraclass coefficient represents the proportion of data variation that can be explained by between-class variability (variability in Der p 1 values between batches). All coefficients were calculated with their 95 % confidence intervals. The level of significance was set at 0.05.

![Figure 1 Linear regression analysis for different pairs of extraction solutions](image-url)
RESULTS

Table 2 shows medians, means, and ranges of Der p 1 mass fractions in dust extracts. Table 3 shows the results of the statistical analysis. Respective Pearson’s correlation coefficients between Der p 1 mass fractions in PBS-T vs. BBS-T, PBS-T vs. ABS-T, and BBS-T vs. ABS-T extracts were 0.967, 0.954, and 0.966 for low exposure range (0.1 μg g⁻¹ to 2 μg g⁻¹) and 0.985, 0.985, and 0.983 for moderate exposure range (2.01 μg g⁻¹ to 7.53 μg g⁻¹). Spearman’s correlation coefficients between Der p 1 mass fractions in the extraction buffers and exposure levels ranged from 0.931 to 1, while intraclass correlation coefficients ranged from 0.927 to 0.993, showing a statistically significant and high agreement in both level ranges. The lowest agreement was observed for Der p 1 measurements in PBS-T and ABS-T extracts (ICC and Pearson’s coefficient of 0.927 and 0.954, respectively) for low exposure range. In the moderate allergen level group, ICC ranged from 0.981 to 0.986 between all three extraction buffers (without significant differences between coefficients), suggesting very good agreements between Der p 1 measurement. Table 3 also shows linear regression coefficients (slope, intercept, and 95 % confidence intervals) for each pair of extraction buffers. Figure 1 shows a very good agreement between Der p 1 measurements between pairs of buffers obtained with linear regression. Each plot shows both the identity line and regression line. The identity line was not included in the 95 % confidence interval for linear regression parameters only in the case of Der p 1 measurements in PBS-T and ABS-T extracts that contained low allergen level. The agreement between all three extraction buffers estimated with ICC was 0.949 (95 % CI: 0.881, 0.98) for low exposure range, 0.983 (95 % CI: 0.952, 0.995) for moderate exposure range, and 0.993 (95 % CI: 0.986, 0.996) overall.

Table 3 Regression analysis (slopes, intercept with 95 % confidence interval) and correlation coefficients (Pearson’s, Spearman’s and intraclass) for Der p 1 mass fractions in PBS-T, PBS-T, and ABS-T extracts

<table>
<thead>
<tr>
<th>Extraction comparison</th>
<th>Der p 1 range / μg g⁻¹</th>
<th>Linear regression</th>
<th>Correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Slope (95 % CI)</td>
<td>Intercept (95 % CI)</td>
</tr>
<tr>
<td>PBS-T vs. BBS-T</td>
<td>0.1 to 2.0</td>
<td>0.906 (0.779, 1.032)</td>
<td>0.043 (-0.07, 0.155)</td>
</tr>
<tr>
<td></td>
<td>2.01 to 7.53</td>
<td>0.988 (0.849, 1.126)</td>
<td>0.23 (-0.338, 0.798)</td>
</tr>
<tr>
<td></td>
<td>overall</td>
<td>1.037 (0.991, 1.084)</td>
<td>-0.018 (-0.137, 0.101)</td>
</tr>
<tr>
<td>PBS-T vs. ABS-T</td>
<td>0.1 to 2.0</td>
<td>0.833 (0.695, 0.971)</td>
<td>0.015 (-0.107, 0.138)</td>
</tr>
<tr>
<td></td>
<td>2.01 to 7.53</td>
<td>0.977 (0.84, 1.115)</td>
<td>0.15 (-0.413, 0.714)</td>
</tr>
<tr>
<td></td>
<td>overall</td>
<td>1.022 (0.973, 1.071)</td>
<td>-0.084 (-0.209, 0.041)</td>
</tr>
<tr>
<td>BBS-T vs. ABS-T</td>
<td>0.1 to 2.0</td>
<td>0.9 (0.772, 1.027)</td>
<td>-0.011 (-0.118, 0.097)</td>
</tr>
<tr>
<td></td>
<td>2.01 to 7.53</td>
<td>0.972 (0.823, 1.121)</td>
<td>-0.01 (-0.646, 0.625)</td>
</tr>
<tr>
<td></td>
<td>overall</td>
<td>0.98 (0.937, 1.023)</td>
<td>-0.057 (-0.171, 0.057)</td>
</tr>
</tbody>
</table>

PBS-T - phosphate buffer-Tween
BBS-T - borate buffer-Tween,
ABS-T - ammonium bicarbonate buffer-Tween,
ICC - intraclass correlation coefficient

All correlation coefficients and regression slopes significantly differed from zero (p<0.001)
DISCUSSION

A number of studies have measured Der p 1 in settled dust worldwide in order to assess exposure risk, especially in children and adults with asthma (4, 13). Monitoring household allergens may play an important role in asthma control (14), but it needs standardised and harmonised protocols for indoor allergen sampling and measurement. Our results show a high correlation and agreement between Der p 1 measurements in PBS-T, BBS-T, and ABS-T at either low or moderate allergen levels. Furthermore, overall ICCs for Der p 1 measurements are high (0.993) for all three buffers. ICC values were slightly lower for the low exposure range than for the moderate because the latter range is wider and involves greater between-class variability and, consequently, a higher intraclass coefficient. Regression analysis showed the slope very close to 1 and small y-axis intercept for each pair of extraction data (Table 3, Figure 1). This reflects high homogeneity of Der p 1 values and excellent agreement between measurements for each pair of extraction buffers (PBS-T, BBS-T, and ABS-T) and may help in standardising the extraction procedure.

Similarly, Martin et al. (15) found no buffer effect on the extraction of Fel d 1 (cat allergen) from dust. Pate et al. (7) also observed that the extraction step was not a significant source of variability. In contrast, Siebers et al. (8) found that borate buffer was superior to PBS and ABS. However, they did not use Tween 20 in that study and their results are not fully comparable with ours. Several investigators reported that adding Tween 20 (as a dispersing and solubilising agent) to the extraction media improved endotoxin detection in dust extracts (16-18). Furthermore, adding Tween 20 to pyrogen-free water has been recommended for endotoxin analysis by the European Committee for Standardization (CEN) (19). However, the influence of Tween 20 on allergen extraction efficiency has not been investigated or its use universally accepted. Therefore, further research should investigate the effect of Tween 20 on extraction efficiency of indoor allergens from dust.

However, variety between laboratories can be great in other operating conditions such as temperature and time of extraction (7). At a lower extraction temperature (4 °C), Sieber et al. (8) reported lower Der p 1 level irrespective of the buffer type. Furthermore, dust sampling and storage may also affect laboratory performance (20, 21). According to Fahibusch et al. (21), storing dust at -20 °C for up to 10 months had no effect on mite allergen levels but Fel d 1 concentration significantly dropped with storage time. However, the freeze-thaw effects on Der p 1 concentrations in dust extracts or dust samples have not yet been investigated.

In 2005, Pate et al. (7) reported the results of the first quality control of common indoor allergen measurements (mite, cockroach, and pets) in residential dust. They found a strong inter-laboratory variability in the levels of all indoor allergens, which pointed to poor standardisation of some steps in allergen measurements. Harmonising protocols for indoor allergen measurement can make results more comparable and lower inter-laboratory variability. Recently, Filep et al. (22) have developed a single standard for eight common indoor allergens. Adding a such control sample with declared allergen levels can improve performance and reduce variability between laboratories.

In this study the efficiency of buffers on Der p 1 extraction from dust samples was compared using correlation coefficients (intraclass, Pearson’s and Spearman’s) and linear regression coefficients. Generally, intraclass correlation is a better indicator of agreement between different measurements than Pearson’s and Spearman’s correlation which may yield misleadingly higher values of agreement in case one extraction solution is constantly giving higher values than the other. Similarly, the linear regression model can produce regression line equal to the identity line even when data points are far from the estimated line. Another drawback of the regression model is the assumption that data points for at least one solution are free of measurement error, which is unrealistic in our case. However, we decided to include linear regression and Pearson’s and Spearman’s correlation coefficients as an addition to intraclass correlation to make possible a comparison with future studies.

CONCLUSION

Our results have shown excellent agreement between Der p 1 measurements regardless of the extraction buffers (PBS-T, BBS-T, and ABS-T) or exposure level. Therefore, all three buffers plus 0.05 % Tween 20 have proved equally efficient in the extraction of Der p 1 from residential dust at room temperature. In order to harmonise extracting procedures, further studies should include extraction from samples with high Der p 1 levels. In addition,
further studies are necessary to find out if the results reported in this study can be generalised for other allergens in reservoir dust samples.

Acknowledgement

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REFERENCES

39. Perfetti L, Ferrari M, Galdi E, Pozzi V, Cottica D, Grignani E, Minoia C, Moscato G. House dust mites (Dep p1, Der f 1), cat (Fel d 1) and cockroach (Bla g 2) allergens in indoor work-places (offices and archives). Sci Total Environ 2004;328:15-21.
**Sažetak**

USPOREDBA PUFERA ZA EKSTRAKCIJU ALERGENA GRINJE Der p 1 IZ PRAŠINE

Der p 1 glavni je alergen grinje *Dermatophagoides pteronyssinus* koji se rutinski određuje u kućnoj prašini. Postupak ekstrakcije Der p 1 iz prašine nije dobro definiran. Cilj je ovoga rada ispitati korelaciju i slaganje između Der p 1 masnih udjela u ekstraktima prašine koji su sadržavali fosfatni (pH 7,4), boratni (pH 8,0) ili amonij-hidrogenkarbonatni (pH 8,0) pufer s dodatkom 0,05 % Tween 20. Dvadeset i osam uzoraka prašine podijeljeno je u tri skupine za ekstrakciju s jednim od tri pufera na sobnoj temperaturi. Maseni udio Der p 1 određen je u ukupno 84 ekstrakta enzim-imunokemijskom metodom (raspon: 0,1 μg g⁻¹ do 7,53 μg g⁻¹). Statističke metode, uključujući i “intraclass” korelaciju, pokazale su visoku korelaciju i slaganje između masnih udjela Der p 1 u svim ekstraktima. Rezultati pokazuju da su sva tri pufera prikladna za ekstrakciju alergena grinje i rutinsko određivanje Der p 1 u prašini.

**KLJUČNE RIJEČI:** alergeni unutarnjih prostora, *Dermatophagoides pteronyssinus*, ekstrakcijski puferi, ELISA, “intraclass” korelacija, sedimentirana prašina

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