The Nearest Neighbor Nuclei Method to Objectify Analysis of Pertussis Toxin-Induced Clustering

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Abstract
The in vivo histamine sensitization test (HIST) has historically been applied to guarantee the safety of acellular pertussis vaccine batches. Non-compliance of batches is primarily associated with the presence of low levels of pertussis toxin (PTx). Because of ethical, standardization and scientific reasons, a variety of alternative in vitro approaches have been studied to replace this lethal HIST. A broadly applied and partially accepted method is the CHO cell clustering test, which is based on the clustered growth pattern of CHO cells when exposed to minute amounts of PTx. One of the major hurdles for global application of CHO clustering test is the manual assessment of the clusters, which is negatively associated with the reproducibility of test outcomes and time consuming. Here, various parameters of CHO cell nuclei were evaluated, in search for a reliable, objective read-out parameter. We demonstrate that the distance between each nucleus and its nearest neighbor (3N method) is the most suitable parameter to assess clustered cell growth. This method detects 2.8 mIU PTx/mL and thereby complies with the requirement set for the sensitivity of the CHO clustering test based on visual reading. In commercial acellular pertussis vaccines spiked with PTx, the method detects 45 mIU/mL PTx, which is substantially lower than the 181-725 mIU/mL PTx detected by visual interpretation. The 3N method thus allows for objective and sensitive assessment of CHO clustering and thereby encourages broad and global implementation of the test as an alternative to the HIST.

1 Introduction
Pertussis toxoid (PTd) is the primary component of acellular pertussis (aP) vaccines and is considered of high importance for vaccine-induced protective immunity (Sato et al., 1984; Black et al., 1988). PTd is generated by inactivation of pertussis toxin (PTx), a toxin that even at low concentrations is responsible for the occasional adverse effects of aP vaccines (Pittman, 1984; Tamura et al., 1982). PTx is an AB5 toxin of which the B-oligomer binds to membrane bound glycoproteins, upon which the toxin enters the cell and is transported to the Golgi and the endoplasmic reticulum. After dissociation from the B-oligomer, the A-subunit is released into the cytosol where it ADP-ribosylates the α-subunit of inhibitory G (Gi)-coupled receptors (Plaut and Carbonetti, 2008; Katada et al., 1983). As a result, the α-subunit can no longer inhibit its target enzyme adenylate cyclase (AC), which converts ATP into cAMP.

Monitoring reversion of the toxoid to its native form as well as assessment of residual PTx has long relied on the lethal murine histamine sensitization test (HIST), in which exposure to PTx increases the sensitivity to histamine by 30-300 times (Kind, 1958; Bergman and Munoz, 1965). This in vivo model, however, causes severe animal suffering, lacks mechanistic understanding, suffers from standardization problems, and accounts for approximately 65,000 mice on a global level each year (Hoonakker et al., 2017).

The CHO cell clustering test is one of several alternatives, some of them still under development (Markey et al., 2019) in which assessment of PTx is based on the PTx-induced clustered growth of CHO cells (Xing et al., 2002; Hewlett et al., 1983). The CHO clustering test is widely applied as a routine test for aP bulk products. Utilization of this test for final aP products has long been hampered by the cytotoxic effect of aluminum salts, an effect which can be circumvented by using semipermeable membranes (Isbrucker et al., 2016). In addition, the requirement to perform the test on the formulated final product has recently been waived by the European Pharmacopeia (Ph. Eur.) (Pharmacopoeia, 2020a). Nevertheless, broad and global implementation...
of the CHO cell clustering test still has to be achieved. One of the major hurdles is the manual reading of the clustering, which is time consuming and associated with variable test outcomes (Markey et al., 2019).

Clustered CHO cells form tightly packed groups resembling a rosette and are typically characterized by a more round morphology as compared to unclustered cells (Hewlett et al., 1983). We hypothesized that these changes may be reflected in the characteristics of the nuclei, which are relatively easy to visualize and measure. We evaluated whether nuclear characteristics were altered as a result of exposure to PTx by analyzing DAPI-stained nuclei. The distance between neighboring nuclei was the only parameter that consistently corresponded with clustering and was therefore selected for further analysis. Using this parameter, we optimized the analysis method and determined the sensitivity to detect PTx alone as well as in the context of two commercial - aP containing - multivalent vaccines. Our findings provide proof-of-principle that measuring the distance between each nucleus and its single most adjacent nucleus, is a sensitive and objective parameter. By objectifying the conventional subjective reading, the 3N method might complement current strategy to waive the HIST and encourage broad application of the CHO cell clustering assay as an in vitro alternative.

2 Materials and methods

2.1 Cell line and culture conditions

Chinese Hamster Ovary (CHO-K1) cells purchased from ECACC (catalogue numbers 85051005, lot 12G006, passage number P+7) were cultured (37°C, 5% CO₂) in F-12 Ham’s (Sigma Aldrich), supplemented with 10% fetal calf serum (v/v), 100 U/ml penicillin, 100 µg/ml streptomycin, 0.3 mg/ml L-glutamine (Gibco). Fetal calf serum was used to adhere to the original clustering protocol (Isbrucker et al., 2016), though replacement of this medium component is envisaged in the future. Cells were passaged every 2-4 days using trypsin-EDTA, and cells of passage number 12-20 were used for the experiments.

2.2 CHO cell clustering assay

Clustering was studied by plating 0.5 mL CHO cells (3 x 10⁴ cells/mL or as indicated) in 24-well plates for 1h at the bench at room temperature and subsequently for 2h in an incubator at 37°C, 5% CO₂. Next, 250 µL medium was removed and replaced by 250 µL of medium or aP vaccine that contained the indicated concentrations of PTx (BRP1 cat#Y0000021; the reference preparation of the European Directorate for the Quality of Medicines & Health Care (EDQM)) for 48h at 37°C, 5% CO₂. One dilution of each was prepared and added to three wells. Cells were fixed with methanol and the nuclei were stained with DAPI (Sigma). CHO cell clustering was visualized using a Leica DMi6000B microscope at a magnitude of 200x. 10x10 images of each well were obtained and stitched using LasX software. 24-well inserts (0.4 µm, Pore Polycarbonate Membrane, Corning) were used when cells were exposed to vaccine. The PTx concentrations in the samples (and not the final concentration) are expressed in units as assigned for the CHO clustering test, one vial containing 50 µg of PTx corresponding to 1360 IU of CHO units (or 7500 IU of HIST units). Two DTAp-IP (Diphtheria, Tetanus, acellular Pertussis and inactivated Polio) vaccines were purchased from Orly Pharma, one being Adacel from Sanofi containing AlPO₃ and the other being Pediarix from GSK containing Al(OH)₃ and AlPO₄. The vaccines were spiked with indicated concentrations of PTx (BRP1) for 1 hour at 4°C on an orbital shaker and centrifuged at 2000 RPM for 10 min at 4°C. The supernatant was discarded to get rid of its cytotoxic components, and the pellet fraction was resuspended in an equal amount of culture medium and stored over night at 4°C until usage.

2.3 Analysis of images

After DAPI staining, the following protocol was used to analyze the images:

— 10x10 images were obtained and stitched
— Using FIJI software, the Gaussian blur filter (tab ANALYZE) was applied with a value of 2.0 (to fill up large gaps in DAPI-stained nuclei)
— Background noise (see tab ANALYZE) was subtracted using a value of 50 (Unick light background)
— Set measurement parameters (tab ANALYZE tick Area, Centroid, Perimeter, Area fraction)
— Choose Type 8-bit (tab IMAGE)
— Subsequently, the threshold is checked and if necessary adapted (tab IMAGE > Adjust > Threshold). Make sure to tick ‘Dark Background’. Document the threshold used (normally between 10 and 20)
— Adjust obtained images to binary images (tab PROCESS > Binary > Make Binary), fill remaining holes (tab PROCESS > Binary > Fill holes), and separate overlapping nuclei (tab PROCESS > Binary > Watershed)
— Analyse the characteristics and positions of the nuclei (tab ANALYZE > Analyze particles and use the following settings: choose Size 250-10.000, Circularity 0.3 – 1, Outlines, and tick Pixels, and Clear results)
— Analyse the distances between nuclei using either of the following plugins: Nnd³ and Simple KNN² (Text S1³). Simple KNN analyses the distance between one or more nuclei neighboring nuclei based on their X and Y coordinates. The

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² https://gist.github.com/lacan/2643f2ce7e33d1bb079a0dfe9f94101, accessed in 2017
³ doi:10.14573/altex.2012171
results of this analysis were used for the figures shown here. The NnD is a simpler script that can only measure the distance between one nucleus and its most adjacent nucleus.

2.4 Statistical analysis
Statistical analysis of nuclear distances was performed with Microsoft R-Open version 3.5.0 (R Core Team, 2018) (script is provided in Text S2²). R is a language and environment for statistical computing. Nuclear distances and the other nuclear parameters (Tab. 1) were log2-transformed to obtain normality. Results are presented as the geometric mean of nuclear distance with corresponding 95% confidence intervals. The log2-transformed nuclear distance for each PTx concentration was compared to medium using a two-sample two-sided t-test, and differences were considered significant when p < 0.05. The Bonferroni method was used for correction of multiple comparisons. Unless mentioned otherwise, the results are expressed as the ratio of the nuclear distances (PTx / Medium) with corresponding 95% confidence intervals (where 1 indicates no difference in nucleus distance and values < 1 indicating clustering). Because a log transformation was applied, the estimated difference in nuclear distance is log2(PTx) – log2(Medium), the antilog of this estimate is a ratio as log a – log b = log a/b.

The number of nuclei required to detect a specified geometric ratio of PTx / Medium was calculated using the following formula:

\[ N = 2 \times (Z_{\alpha/2} + Z_{\beta})^2 \times \left(\frac{SD}{ES}\right)^2 \]

Where: \( N \) = required number of nuclei, \( Z_{\alpha/2} \) = Critical value for type I error = 1.96 (\( \alpha = 0.05 \)), \( Z_{\beta} \) = Critical value for type II error = 2.33 (1 – \( b = 0.99 \)), \( ES \) = effect size, expressed as log2(ratio), \( SD \) = Standard deviation estimate based on three experiments.

3 Results
3.1 Selection of parameters to assess clustering
CHO cell nuclei were stained with DAPI (Fig. 1), and acquired images were converted to binary files. These files were used to evaluate nuclear size, morphology, and the distance between adjacent nuclei. While non-exposed cells and nuclei were randomly distributed (Fig. 1A), the distribution of both PTx exposed cells as well as their DAPI-stained nuclei was characterized by a clustered pattern (Fig. 1B). Exposure to increasing PTx concentrations was associated with a decrease in overall cell numbers, but the high variability impeded the detection of significant effects (Tab. 1). The morphology of the nuclei (roundness and aspect ratio) was slightly, but significantly, altered upon exposure to 2.8, 11 and 181 mIU/mL PTx (Tab. 1), while 45 mIU/mL PTx had no effect. In addition, nuclear circularity was slightly reduced when cells were exposed to PTx down to a concentration of 11 mIU/mL (Tab. 1). Assessment of nuclear sizes (area) demonstrate that these were significantly increased when cells had been exposed to PTx with a detection level of 2.8 mIU/mL PTx, but there was no clear dose-response effect (Tab. 1) and the distribution was not normal (Fig. 1C). Most importantly, the distance between one nucleus and its nearest neighboring nucleus was significantly reduced when cells were exposed to PTx down to a concentration of 2.8 mIU/mL (Tab. 1). The percentage change in this distance increased with increasing PTx concentrations (Tab. 1) and was normally distributed, though slightly skewed towards the right (Fig. 1D). Consequently, the distance between nearest neighboring nuclei was selected as the most promising parameter.

3.2 The effect of combining triplicate measures versus individual measures
The analysis of the PTx reference and vaccine bulk samples in monograph 2.6.33 of the Ph. Eur. on ‘residual pertussis toxin’ is based on at least dual measurements. In Figure 1 and Table 1, statistical analysis was performed by combining all nuclei in three wells (i.e., triplicate measures) to ensure optimal statistical resolution. Since analysis of the nuclei in one well (i.e., single measures) might be more practical, statistical analyses were performed either by combined analysis of the triplicates or by performing the analysis on single measures. In both the triplicate (Fig. 2A, Fig. S1A,B) as well as individual analysis (Fig. 2B, Fig. S1C,D), the effect of PTx was either pronounced (11-725 mIU/mL) or slight (2.8 mIU/mL). The individual analysis is associated with substantial variation between triplicates, which is compensated by combining the statistical analyses of triplicates. We therefore considered combined analysis as optimal for PTx induced clustering.

3.3 Nuclear numbers required for reliable analyses
In addition, the minimal number of cells required for reliable analysis was evaluated statistically (Fig. 2C) and experimentally (Fig. S2²). Figure 2C shows that the total number of nuclei required for reliable analysis is 302 nuclei for PTx in the concentration range of 11-725 IU/mL and 6294 nuclei for a concentration of 2.8 mIU/mL. The average number of nuclei in each binary image is approximately 3500, explaining the necessity to combine the analysis of triplicates for detection of 2.8 mIU/ml PTx. When we experimentally halved cell numbers (Fig. S2A²) the ratios PTx/medium were slightly enhanced (average ratio PTx/medium was 0.90 instead of 0.86), but the lowest detectable PTx concentration remained 2.8 mIU/mL. By contrast, experimentally increasing cell numbers (Fig. S2C²) did not substantially influence the ratio PTx/medium but slightly reduced the sensitivity to detect PTx (11 mIU/mL PTx instead of 2.8 mIU/mL PTx). These minimal effects of cell densities underline the robustness of the assay.
**Fig. 1: The effect of PTx on the nuclear distribution of CHO cells**

CHO cells were exposed to medium (A) or 181 mIU/mL PTx (B) for 48 hours and nuclei were visualized with DAPI (shown in blue/purple). Fluorescence images were converted to binary images. Shown are examples of one overlay (fluorescence and light microscopical, 1x image) of each exposure condition and corresponding stitched binary images (composed of 10x10 images). Based on the analysis of all nuclei in three wells, the area of the nuclei (C) and the distance between the nearest neighboring nuclei (D) were determined. Shown is one out of two representative experiments. The lines indicate the geometric mean of three wells for each condition.

**Fig. 2: The distance between nearest neighboring nuclei using triplicates or single measurements (experiment 1)**

CHO cells were exposed to 0, 2.8, 11, 45, 181 and 725 mIU/mL PTx for 48 hours and nuclei were visualized with DAPI. Fluorescence images were acquired, stitched (10x10 images) and converted to binary images. Shown is the ratio of the geometric mean distance between nuclei for the indicated PTx concentration divided by the geometric mean distance between nuclei for medium, and the confidence interval for each concentration of PTx. Analyses was based on all nuclei in three wells (A) or on the nuclei in the individual wells (B). One out of three representative experiments is shown, the other two experiments are shown in Figure S1. (C) The number of nuclei required for reliable analysis for each ratio of PTx/medium, based on three experiments. Highlighted in grey (A, C) is the range of ratios for 11-725 IU/mL PTx and in blue the ratio for 2.8 IU/mL PTx.
Tab. 1: CHO cell clustering and the associated changes in DAPI stained nuclei
Number: the number of nuclei detected in the analysed field. Roundness (Round): 4 × ((Area)/(π×(Major axis)²)). Aspect ratio (AR): The aspect ratio of the particle's fitted ellipse, here (Major Axis)/(Minor Axis). Circularity (Circ.): 4π × ((Area)/(Perimeter)²) with a value of 1.0 indicating a perfect circle. As the value approaches 0.0, it indicates an increasingly elongated shape. Area: Area of selection (= nucleus) in square pixels. Distance: distance between two closest adjacent selections (= nucleus). Shown is the geometric mean of the lowest concentration detected based on the point that is significantly different from 0 IU/ml PTx; vs., versus; AU, arbitrary units; IU, international units.

<table>
<thead>
<tr>
<th>PTx (mIU/mL)</th>
<th>Growth pattern</th>
<th>Number</th>
<th>Round (AU)</th>
<th>AR (AU)</th>
<th>Circularity (AU)</th>
<th>Area (AU)</th>
<th>Distance (AU)</th>
</tr>
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<tbody>
<tr>
<td>181</td>
<td>Clustered</td>
<td>5884</td>
<td>3%</td>
<td>0.662</td>
<td>1.51 (1.51 - 1.52)</td>
<td>0.812 (0.811 - 0.813)</td>
<td>0.207 (0.209)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(4790 - 7228)</td>
<td></td>
<td>(0.660 - 0.664)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>45</td>
<td>Clustered</td>
<td>5227</td>
<td>-8%</td>
<td>0.679</td>
<td>1.47 (1.47 - 1.48)</td>
<td>0.822 (0.821 - 0.823)</td>
<td>0.219 (0.220)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(3919 - 6971)</td>
<td></td>
<td>(0.677 - 0.681)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>Clustered</td>
<td>6563</td>
<td>15%</td>
<td>0.669</td>
<td>1.50 (1.49 - 1.50)</td>
<td>0.832 (0.831 - 0.832)</td>
<td>0.217 (0.218)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(5940 - 7251)</td>
<td></td>
<td>(0.667 - 0.671)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Clustered</td>
<td>6548</td>
<td>15%</td>
<td>0.672</td>
<td>1.49 (1.49 - 1.49)</td>
<td>0.841 (0.840 - 0.842)</td>
<td>0.225 (0.226)</td>
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<tr>
<td></td>
<td></td>
<td>(5186 - 8269)</td>
<td></td>
<td>(0.670 - 0.674)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>Normal</td>
<td>5686</td>
<td>0%</td>
<td>0.675</td>
<td>1.48 (1.48 - 1.49)</td>
<td>0.844 (0.843 - 0.844)</td>
<td>0.206 (0.207)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(4539 - 7122)</td>
<td></td>
<td>(0.673 - 0.677)</td>
<td></td>
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<td></td>
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</table>

Fig. 3: Statistical analysis of the distance to one, two, three or four nearest neighboring nuclei
CHO cells were exposed to 0, 0.7, 2.8, 11, 45, 181, 725 and 2901mIU/mL PTx for 48 hours and nuclei were visualized with DAPI. Fluorescent images were acquired, stitched (10x10 images) and converted to binary images. Shown is the ratio of the geometric mean distance between nuclei for the indicated PTx concentration divided by the geometric mean distance between nuclei for medium, for one (A), two (B), three (C) or four (D) nearest neighboring nuclei. Results of three independent experiments are shown (Exp 1, 2 and 3) and the confidence interval for each concentration of PTx. The black lines indicate a ratio of 1. AU: arbitrary units.

3.4 Analysis of one to four nearest neighboring nuclei
Although the distance between one nucleus and its closest adjacent neighbor nucleus already enables detection of levels as low as 2.8 mIU/mL PTx, analysis of the distance of each nucleus to two or more adjacent nuclei might improve the sensitivity of the assay (Fig. 3). By analyzing the distance of each nucleus to its single nearest neighbor, the lowest concentration detected was between 0.7 and 11 mIU/mL PTx, whereas the lowest concentration detected based on the analysis of the distance to its two, three or four nearest nuclei was between 2.8 and 11mIU/mL PTx (Fig. 3). Repetitions of this experiment which included several concentrations within the range of the lowest concentrations detected, revealed that analysis of an increasing number of nuclei concurred with an increase in variation towards the point that PTx effects were no longer significant for four nuclei in one out of three experiments (Fig. S3). In the other two experiments (Fig. S3), the ratio between cells treated with and without PTx was similar for one and for two nuclei but was reduced when higher numbers of neighboring nuclei were analyzed. Therefore, analysis of one or two nuclei is considered optimal. For practical reasons, one neighboring nucleus was analyzed in the subsequent experiments.
Experiment 1

PTx (mIU/mL) | 0    | 2.8   | 3.8   | 5.7   | 11   | 45    | 161   | 725   \\
---|------|------|------|------|------|------|------|------\
Ratio (low - upp) | -    | -    | 0.97 | (0.99 - 0.95) | 0.95 | (0.97 - 0.94) | 0.92 | (0.93 - 0.91) | 0.92 | (0.94 - 0.91) | 0.87 | (0.89 - 0.85) | 0.88 | (0.89 - 0.86) | 0.87 |
Experiment 2

PTx (mIU/mL) | 0    | 2.8   | 3.8   | 5.7   | 11   | 45    | 161   | 725   \\
---|------|------|------|------|------|------|------|------\
Ratio (low - upp) | -    | -    | 0.96 | (0.95 - 0.97) | 0.96 | (0.95 - 0.97) | 0.97 | (0.96 - 0.98) | 0.86 | (0.86 - 0.87) | 0.82 | (0.81 - 0.83) | 0.82 | (0.81 - 0.83) | 0.82 |
Experiment 3

PTx (mIU/mL) | 0    | 2.8   | 3.8   | 5.7   | 11   | 45    | 161   | 725   \\
---|------|------|------|------|------|------|------|------\
Ratio (low - upp) | -    | -    | 0.95 | (0.95 - 0.96) | 0.96 | (0.95 - 0.96) | 0.92 | (0.92 - 0.93) | 0.92 | (0.91 - 0.92) | 0.83 | (0.82 - 0.84) | 0.82 | (0.81 - 0.83) | 0.81 |

Fig. 4: CHO cell clustering in response to PTx of LIST Biological

CHO cells were exposed to 0 and 0.005-4.0 ng/mL PTx (LIST Biological) for 48 hours and nuclei were visualized with DAPI. Fluorescent images were acquired, stitched (10x10) and converted to binary images. Shown is the ratio of the geometric mean distance between nuclei for the indicated PTx concentration divided by the geometric mean distance between nuclei for medium, and the confidence interval for each concentration of PTx. The black line indicates a ratio of 1. AU: arbitrary units.

The three experiments provide an indication for the reproducibility depicted in more detail in Table 2. There is some variability between the three experiments regarding the absolute distance and a slight variation for the ratios PTx/med. However, in all three experiments the 3N method consistently detects a significant effect of 2.8-725 mIU/mL PTx. Nevertheless, formal assessment of the assays sensitivity and reproducibility need to be addressed in a (pre)validation study.

3.5 A second source of PTx

Although the European PTx standard (BRP1) is commonly applied within and outside Europe, PTx is produced by many others, and the biological activities of these preparation are highly variable and do not necessarily correspond with their protein concentration. Therefore, compatibility of the 3N methods with other sources of PTx is of importance. One commonly used source is the PTx produced by LIST Biological, for which clustering based on visual reading has been detected with concentrations as low as 0.025-0.1 ng PTx/mL (Wagner et al., 2017). Using the 3N method, clustering could be detected with concentrations as low as 0.005 ng PTx/mL PTx (Fig. 4).

3.6 Analysis of PTx in aP vaccines

Although the Ph. Eur. waived the requirement to perform the in vivo HIST for testing of residual PTx and reversion to toxicity of all PTd and final products, the test is still stipulated for these vaccines in other regions of the world. Compliance of the method proposed in this study with formulated aP vaccines would therefore greatly facilitate broad and global application. Preliminary experiments showed that aP vaccine supernatant was cytotoxic to CHO cells and made a membrane necessary to prevent detrimental
Fig. 5: The effect of PTx-spiked aP vaccines on the distance between each nucleus and its most adjacent nucleus

Two DTaP-IP vaccines (Adacel from Sanofi containing AlPO₄ (A) and Pediarix from GSK containing Al(OH) and AlPO₄(B)) were spiked with 0, 11, 23, 45, 91 and 181 mIU/mL PTx. CHO cells were exposed to these vaccine samples or medium for 48 hours and nuclei were visualized with DAPI. Fluorescence microscopical images were acquired, stitched (10x10) and converted to binary images (using ImageJ). Shown is the geometric mean distance between one nucleus to its nearest neighbor and the confidence interval for each concentration of PTx for one experiment. The red lines indicate the confidence interval for 0 IU/mL PTx (in vaccine). AU: arbitrary units.

effects of the adjuvant (data not shown). Consequently semipermeable membranes were applied and used to evaluate the pellet fraction of two commercial aP vaccines, Adacel containing AlPO₄ and Pediarix containing Al(OH) and AlPO₄. The plain aP vaccines are not inert and they reduced cellular confluency and slightly increased the distance between the nuclei as compared to cells grown in medium alone (Fig. 5A, B), which might be the results of the adjuvant or adsorbed vaccine components. PTx in a concentration range of 11–181 and 45–181 mIU/mL spiked into Adacel and Pediarix respectively, significantly reduced the distance between one nucleus and its most adjacent neighbor nucleus as compared to unspiked vaccine. The 3N method was therefore capable of detecting lower concentrations of PTx than included in the collaborative study using visual inspection of clustering (Isbrucker et al., 2016). Depending on the laboratory and the vaccine, the lowest concentrations detected in four aP vaccines was either 181, 363 or 725 mIU PTx/mL.

4 Discussion

The current reading of the CHO clustering assay is based on visual assessment of the cellular growth pattern, which is labor intensive and requires specific training. Due to its subjective nature, the method is inherently prone to intra- and interlaboratory variation (Xing et al., 2010), although recent studies showed significant improvement by applying an appropriate reference and a standardized protocol (Isbrucker et al., 2016; Markey et al., 2018). Despite attempts by different laboratories, a reliable and objective parameter for clustering has not been identified yet. When clustered, the CHO cell morphology changes, the cells become more packed and form islets (Fig. 1). As nuclei are relatively easy to stain, we evaluated whether clustering was associated with changes in nuclear numbers, size, shape, and in the distance between nuclei. Our analyses showed that the distance between nearest neighboring nuclei (3N) was the only parameter that consistently corresponded with clustering. Statistical evaluation of various factors demonstrates that analysis of the distance of each nucleus to its single most adjacent nucleus and combining the counts of cells in three wells is the best procedure to determine CHO clustering in an objective manner.

In principle, the CHO clustering test is a limit test for detection of PTx in intermediate aP vaccine products, although evaluation of dilution ranges of products also allow for the approximative assignment of PTx concentrations (Pharmacopoeia, 2020b). Using our 3N method, we show that the distance between nuclei of CHO cells is linear between 5.7 and 45 mIU/mL PTx (Fig. S1³), although the range of this linear part varies somewhat between experiments (Tab. 2). We therefore recommend assessment of nuclear distances as a qualitative assay. For such qualitative assays, the ICH requires the evaluation of the detection limit and the specificity (ICH, 2005). Using the European reference PTx standard (BRP1), the detection limit for the CHO clustering test based on visual examination is 5 mIU/mL (CHO units) (Pharmacopoeia, 2020b), while lowest concentration detected with the 3N method is 2.8 mIU/mL PTx. The source of PTx commonly used in the US is LIST Biological, for which the CHO clustering limit is 0.025–0.1 ng/mL PTx (Wagner et al., 2017), while it is 0.019 ng/mL PTx using the 3N method. The European requirement for the sensitivity of the CHO clustering provides an important and central point of focus for the assay sensitivity using purified PTx. For final product testing, there is consensus that an in vitro alternative should have a sensitivity at least equivalent to the HIST (Arciniega et al., 2016; Wagner et al., 2016), which nevertheless poses a challenge due to the diverse sensitivities of regional
variants of this in vivo test (Arciniega et al., 2016). Using the European Pharmacopeia method, the HIST has a detection limit of 1–2 IU (HIST units corresponding to 181–363 mIU CHO cell units) PTx in aP vaccines (Xing et al., 2010; Bache et al., 2012). Here, experiments with two commercial aP vaccines spiked with purified PTx demonstrate that the lowest detectable concentrations are as low as 11 and 45 IU/mL PTx for Adacel and Pediarix respectively by measuring the distance between nuclei. This is well below the sensitivity of the HIST and below the lowest PTx concentration included in a collaborative study based on visual reading of CHO clusters (Isbrucker et al., 2016). Some regulatory agencies require testing for residual PTx as well as reversion of PTd to its toxic forms (Hoonakker et al., 2017), and a follow up validation study should therefore determine the assay’s detection limit for both purposes using the 3N method.

The other criterion of the ICH, specificity, is described as "the ability to assess unequivocally the analyte in the presence of components which may be expected to be present" (ICH, 2005). The two vaccines used in this study were produced by Sanofi and GSK and differ in their matrix and adjuvant composition, factors known to affect CHO cell clustering (Isbrucker et al., 2016). Here, we show that the distance between nuclei increases as a result of exposure to these commercial aP vaccines (Fig. 4). This is most likely the result from a decrease in cellular confluency caused by the adjuvant fraction or minute amounts of the diluent fraction that might be slightly toxic to the CHO cells, even when separated from the cells with a semipermeable membrane (Isbrucker et al., 2016). These slight but significant effects can be compensated for by normalization using a safe batch of the same composition as the vaccine under study in each experiment, according to the principles of the consistency approach (De Mattia et al., 2011). The compliance of the 3N method with both ICH criteria underscores the suitability of the method to assess clustered growth and thereby contributes to application of the CHO clustering test for final product assessment. Nevertheless, further research should include in-house validation studies, evaluating the method’s reproducibility and sensitivity by using blinded spiked samples. In that respect, the 3N method could be applied side-by-side with the conventional reading method.

We here show that the 3N method is valid for assessing clustered growth of CHO cells. Scanning of the plates, preparation of the binary pictures, and the statistical analysis can and should be automated to reduce hands-on-time up to 2–4 times. Our study provides proof-of-principle that measuring the distance between nearest neighboring nuclei is a sensitive and objective read-out for CHO cell clustering exposed to the different PTx preparations used in this study. This indicates that the method may be applied for bulk product testing within Europe, although it should be subjected to further in-house validation before application. Furthermore, we demonstrate the validity of this method for testing of final vaccine products, which could endorse implementation of the CHO clustering test as an alternative for the HIST on a global level.

References


Conflict of interest statement
The authors declare no financial or commercial conflicts of interest.

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