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ARTICULO ORIGINAL

### Di- and triploid erythrocyte identification by multi-parameter image analysis: A new method for the quantification of triploidization rates in rainbow trout (*Oncorhynchus mykiss*)

Identificación de di- y triploidización por análisis multiparamétrico de imágenes: Un nuevo método para la cuantificación de la tasa de triploidización en trucha arcoiris (*Oncorhynchus mykiss*)

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#### RESUMEN

La creciente competencia internacional ha forzado a la industria del salm n a la incorporaci n de t cnicas innovadoras. El cultivo de hembras triploides tiene m ltiples ventajas sobre poblaciones diploides. En la actualidad, no existe un m todo simple, exacto y de bajo riesgo para la cuantificaci n de tasas de triploidizaci n. En este trabajo presentamos un m todo que

multiparamétrico de imágenes, denominándolo como microscopía morfológica cuantitativa (QMM). Se utilizó citometría de flujo (FC) como un método de referencia para determinar el contenido de ADN en eritrocitos diploides y triploides extraídos de truchas arco iris inmaduras (*Oncorhynchus mykiss*). Además, se aplicó microscopía de fluorescencia cuantitativa (QFM), usando los marcadores de ADN: 4,6-diamidina-2-fenilindol (DAPI), Yoduro de Propidio (PI), y Naranja de Acridina (AO). Nuestros datos muestran que QMM posee una capacidad discriminante comparable o incluso superior a FC o QFM. El método desarrollado ofrece una nueva perspectiva para clasificar objetos microscópicos, con muchas posibles aplicaciones.

*Key words:* triploidization, cytometry, microscopy, image processing.

*Palabras clave:* triploidización, citometría, microscopía, imagen procesada.

## ABSTRACT

Growing international competition is forcing salmon farmers to incorporate innovative techniques into the production process. The use of triploid, all-female breeding populations offers multiple advantages over diploid populations. Currently, an exact, simple, and non-hazardous method for the quantification of diploid- and triploid salmon erythrocytes does not exist. We present a method that combines a standard microscopic bright field technique (contrast staining with GIEMSA) with multi-parameter image analysis and termed it quantitative morphologic microscopy (QMM). We used flow cytometry (FC) as the reference method to determine the DNA content of di- and triploid erythrocytes from immature rainbow trout (*Oncorhynchus mykiss*). Additionally, we applied quantitative fluorescence microscopy (QFM), using the DNA stains 4',6-diamidino-2-phenylindole (DAPI), propidium iodide (PI), and acridine orange (AO). Our data show that QMM possess comparable or even superior discriminating capacities than FC or QFM. The developed method opens a perspective for the classification of microscopic objects with many possible applications.

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## INTRODUCTION

The economic situation has forced the salmon industry to maximize production rates, improve fillet quality, and minimize breeding costs. The use of all-female, triploid lines provides advantages in comparison to non-sterile diploid females, male, or sexually mixed populations (Utter *et al* 1983, Wlasow *et al* 2004, Friars *et al* 2001). Female triploid salmon do not develop ovaries, nor do they express the characteristic phenotypic changes of the sexual maturation of the diploid species (Happe *et al* 1988). Methods for triploidization include the application of high pressure or hyperthermia after the fertilisation procedures. Presently, high pressure treatment is preferred because it yields better triploidization and surviving rates. Since young di- and triploid fishes cannot be distinguished by their phenotypes, triploidization rates are determined on a cellular basis with different protocols (Thititananukij *et al* 1996, Boron 1994, Child and Watkins 1994, Wattendorff 1986, Ewing and Scalet 1991, Cozier and Moffett 1989, Al-Sabti 1995, Thomas and Morrison 1995). Due to its precision, the reference method for the quantification of the nuclear DNA content is based on fluorescent staining of the cellular chromatin in combination with flow cytometry (FC) (Utter *et al* 1983). The main disadvantages of FC are high costs and the handling of hazardous DNA binding fluorescent dyes, such as acridine orange (AO), propidium iodide (PI), or 4',6-diamidino-2-phenylindole (DAPI). So far, an easy, precise, and non-hazardous method for the quantification of triploidization rates has not been developed. Applied methods are based on the fish to fish determination of triploidy in erythrocytes by trained specialists who classify up to 100 samples in order to obtain statistically significant results. In this work we present a microscopic bright field technique for the classification of di- and triploide erythrocytes of *O. mykiss*. This technique takes advantage of image processing routines in combination with multiple component analysis. The results obtained with quantitative morphologic microscopy (QMM) or quantitative fluorescent microscopy (QFM) are comparable or even superior to the FC-data. QMM substitutes fluorescence parameters for pure morphologic features of the erythrocyte nuclei which can be contrasted by simple GIEMSA staining procedures. QMM offers a precise, non-hazardous, inexpensive, and relatively simple method for classification of di- and triploid erythrocytes of *O.*

## MATERIAL AND METHODS

*Triploidization method & collection of blood samples:* Triploidization of rainbow trout (*Oncorhynchus mikiss*, Smith and Stearly 1989) was induced by hyperthermia (28°C for 10 min), 40 min after fertilisation. Blood samples (1 ml) were collected from immature fishes (350 ± 50 g) by piercing the caudal vein. Heparin (Sigma) was added to inhibit blood coagulation.

*Discrimination of di- and triploid erythrocytes by flow cytometry (FC):* Following the method described by Darzynkiewicz *et al* (1979), 100 ml blood suspension (10<sup>5</sup> erythrocytes) was mixed with 200 ml of acid detergent (0.1 % Triton X-100, 0.08 N HCl, 150 mM NaCl). After 30 s, 600 ml of the acridine orange (AO) staining solution was added (100 ml contained 1.2 mg AO (Sigma), 29 mg ethylene-diamine-tetra-acetic acid (EDTA), 150 mM NaCl, 0.1 M C<sub>6</sub>H<sub>8</sub>O<sub>7</sub>H<sub>2</sub> O, and 0.2 M Na<sub>2</sub>HPO<sub>4</sub>). All steps were performed in a 0°C ice water bath. After 10 min, the DNA content of 5000 cell nuclei was determined by FC as described by Ojeda *et al* (1992). Excitation of DNA-bound AO was performed with a 50 mW laser at 488 nm, while the emission intensity was recorded at 515-575 nm for each individual cell. Additionally, side scattering (90°) and forward scattering were recorded for each cell. The setup of the FC, designed at the Institute of Physics of the Universidad Austral de Chile by Flavio Ojeda, is described in Ojeda *et al* (1992).

*Discrimination of di- and triploid erythrocytes by quantitative fluorescence microscopy (QFM):* Blood suspensions (5 ml) were pipetted onto different microscopic slides. Erythrocyte nuclei were stained with AO staining solution (see above), propidium iodide (PI) (Molecular Probes, Eugene, OR, USA) staining solution (50 mg/ml), or 4',6-diamidino-2-phenylindole dihydrochloride (DAPI) staining solution (50 mg/ml). For AO and PI staining, 5 ml of acid detergent was added to the slides in order to permeabilize the plasma membranes of the erythrocytes. After 5 min, 5 ml of the corresponding staining solutions were added. For DAPI staining, 5 ml staining solution was added directly to the erythrocytes, since the small dye diffuses freely into the chromosomal region. After an incubation time of 10 min, samples were covered with slide covers and observed on a Leica epifluorescence microscope DM LB, equipped with a type A filter set and a CS 50 W-4/L2 lamp for the fluorescent and the bright field mode. Gray scale images (8-bit, 701 × 480 pixels) were recorded using a CCD camera (CE, Model OS70D HR, Taiwan) and GrabIT frame grabber soft- and hardware (AIMS Lab., USA).

*Discrimination of di- and triploid erythrocytes using bright field quantitative morphological microscopy (QMM):* Small aliquots of blood samples were spread out on microscopic slides and dried at room temperature. Samples were covered with May Grünwalds solution (SIGMA) for 1 min, covered with aq<sub>dest</sub> for 3 min, washed with aq<sub>dest</sub>, covered with GIEMSA solution (SIGMA, 50 ml/ml aq) for 15 min, and washed with aq<sub>dest</sub>. Finally, samples were covered with cover slides and images were recorded in the bright field mode with the microscopic setup described above.

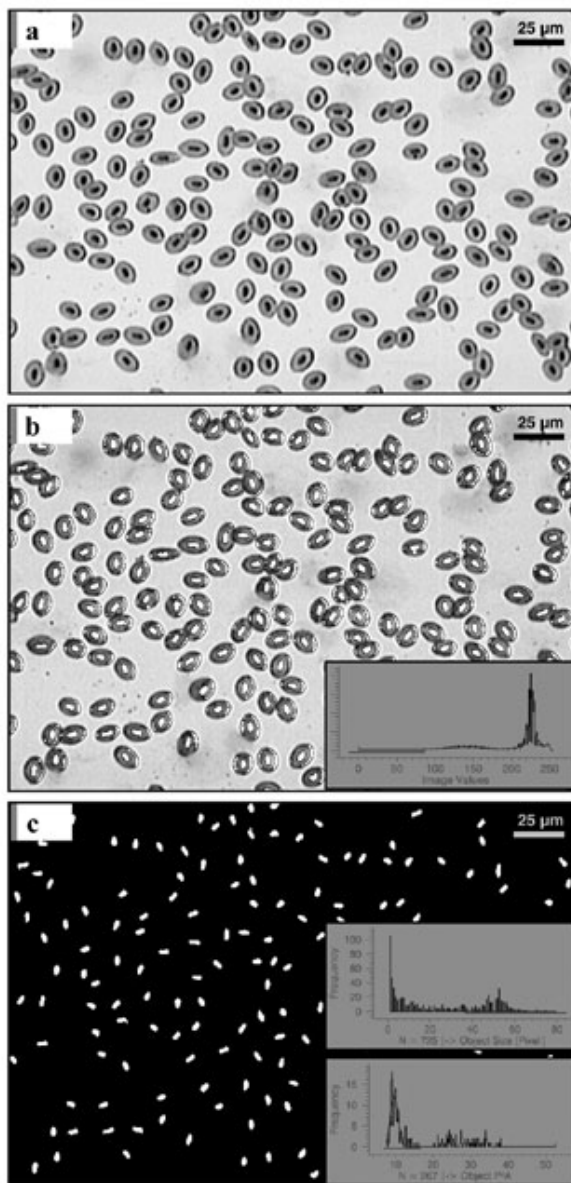
*Segmentation and parameterization of erythrocyte nuclei:* Interactive routines for cellular image analysis and multivariate data processing (SCIAN) were written in IDL<sup>®</sup> 5.4 (Interactive Data Language, Research Systems, CO, USA). SCIAN ([www.scian.cl](http://www.scian.cl)) contains a library of different filters which allow the segmentation of complex structures (Fanani *et al* 2002, Härtel *et al* 2005), cells (Jessel *et al* 2002, Härtel *et al* 2003), or higher biological samples (Alvarez *et al* 2004/2005). For fluorescent images, threshold segmentation in the intensity-histogram yielded excellent results. For bright field images, either threshold segmentation in the intensity-histogram ([figure 1](#)) or a modified scaling index method (SIM) was applied. SIM was originally introduced by R ath and Morfill (1997) as a pixel-wise non linear transformation. We introduced the weighting vector  $p$  in [2] and [3] in order to optimize the segmentation results:

$$\alpha(x_i, y_i | r_1, r_2) = \frac{\log[N(r_2)] - \log[N(r_1)]}{\log(r_2) - \log(r_1)} \quad [1]$$

$$N(x_i, y_i | r_1) = \sum_{x,y} \Theta(r_1 - |\bar{p}_i - \bar{p}|) \cdot \left( \frac{r_1}{|\bar{p}_i - \bar{p}|} \right) \quad [2]$$

$$N(x_i, y_i | r_2) = \begin{cases} \sum_{x,y} \Theta(r_1 - |\bar{p}_i - \bar{p}|) \cdot \left( \frac{r_1}{|\bar{p}_i - \bar{p}|} \right) & \cdot |\bar{p}_i - \bar{p}| \leq r_1 \\ \sum_{x,y} \Theta(r_2 - |\bar{p}_i - \bar{p}|) \cdot \left( \frac{r_2}{|\bar{p}_i - \bar{p}|} \right) & \cdot |\bar{p}_i - \bar{p}| > r_1 \end{cases} \quad [3]$$

Vector  $\bar{p}$  is defined at each picture position  $(x,y)$  as  $\bar{p} = [x, y, I(x,y)] = [x, y, I(x,y)]$ .  $Q$  is the Heaviside Function and  $||$  the Euclidean Norm. We defined  $r_1$  and  $r_2$  with 4 parameters  $[r'_1 < r'_2, f_1 \& f_2]$ , yielding  $(r_{1/2})^2 = (r'_{1/2})^2 + (f_{1/2})^2$ .  $S$  is defined for  $(x_i - x)^2 + (y_i - y)^2 \& r^2$ . Nuclei are segmented by selecting a threshold value in the transformed picture  $a(x,y)$ . Consecutive erosion and dilation operators correct remaining segmentation defects. Debris and nuclei which touch the image borders are removed by morphological filters which identify the pure nuclei population ([figure 1b/c](#)). For each segmented nucleus, morphological and intensity dependent parameters were calculated ([table 1](#)). Ten pictures containing 1500 to 4500 nuclei were used for QFM and QMM.



**Figure 1.** Segmentation and morphologic filtering of GIEMSA-stained triploid erythrocyte nuclei of *O. mykiss*. [a] Original image as recorded by the CCD camera (8 bits, 634 × 428 pixels). [b] White regions are segmented in a first segmentation step. The inset shows the intensity-histogram of the original image. The selected threshold-interval  $I(x,y) \in [0, 88]$  is marked by a bar below the plot. [c] White regions show triploid erythrocyte nuclei after the application of two morphologic filters. Insets show the frequency histograms of the morphologic parameter size (upper histogram) and  $P^2/A$  (lower histogram). Selected threshold-intervals are marked by bars below the plots. Triploid nuclei cover an area of 36-80 pixels and provide characteristic  $P^2/A$ -values between 8-17. Bars in the upper right corner of the images scale 25 mm.

Segmentación y filtrado morfológico de núcleos de eritrocitos triploides de *O. mykiss*. [a] Imagen original digital (8 bits, 634x428 píxeles). [b] Estructuras segmentadas marcadas en blanco después del primer paso de segmentación. El histograma interior muestra las frecuencias de las intensidades de la imagen original. El umbral seleccionado  $I(x,y) \in [0, 88]$  para la segmentación está marcada con una barra. [c] Estructuras marcadas en blanco muestran los núcleos de eritrocitos después de la aplicación de dos filtros morfológicos. Los histogramas interiores muestran el filtro del tamaño del parámetro morfológico (recuadro superior) y  $P^2/A$  (recuadro inferior). Los histogramas contienen los umbrales usados para filtrar estructuras perinucleares.

**Table 1.** Morphological, texture, and fluorescence dependent parameters available in SCIAN for the classification of segmented objects in digital images (figure 1c).

Parámetros dependientes de morfología, textura y fluorescencia dentro de SCIAN para la clasificación de objetos segmentados en imágenes digitales (figura 1c).

Morphology, Texture, and Fluorescence Dependent Parameters:	
Size (A).	Number of pixels identifying each erythrocyte nuclei.
Perimeter (P) of the trajectory of the object boundary.	Boundary pixels (4-connected contour, Young <i>et al</i> 1997) of the nuclei were translated into chain code according to Freeman (1979). Boundary length was then determined according to Vossepoel (1982).
$P^2/A$ .	Dimensionless, zoom, and rotation invariant circularity parameter.
Minimum border distance (D) and $\underline{D}$ .	$D = \sum_{i=1}^A  d_{min} _i$ and $\underline{D} = \frac{1}{A} \cdot \sum_{i=1}^A  d_{min} _i$ , $ d_{min} _i$ is the minimum Euclidean Distance between each pixel of an object and its nearest boundary pixel (Danielson, 1978).
Boundary energy ( $E_C$ ). Normalized boundary energy ( $\underline{E}_C$ ).	$E_C = \oint  K(p) ^2 dp$ and $\underline{E}_C = \frac{1}{P} \oint  K(p) ^2 dp$ were introduced by Young <i>et al.</i> (1974). $K(p) = 1/r(p)$ is the curvature, and $r(p)$ is the radius of the curvature of the boundary trajectory p.
Number of turning points ( $2^{nd}_d$ ) and $2^{nd}_d/P$ .	Turning points of the boundary trajectory can be identified when $K(p) = 0$ and the circle touching the trajectory tangentially, switches the side of the boundary. $2^{nd}_d$ is the number of turning points for each boundary trajectory and $2^{nd}_d/P$ is the ratio $2^{nd}_d/P$ .
Fluorescence intensity (I) and fluorescence density (I').	$I = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} I(x', y') dx' dy'$ , and $I' = \frac{1}{A} \cdot \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} I(x', y') dx' dy'$ .
Normalised translation and rotation invariant moments ( $\mu'_{jk}$ ) and ( $ \mu'_{jk}$ ).	$\mu'_{jk} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} (x' - \bar{x})^j \cdot (y' - \bar{y})^k \cdot f(x', y') dx' dy'$ . Were calculated for each nucleus after transformation on its principal axis and in relation to its centre of gravity (Castleman 1996). For morphological invariant moments $\mu'$ of order jk, $f(x', y')$ is set to 1. For texture or fluorescence invariant moments $ \mu'$ , $f(x', y')$ is set to the intensity value $I(x', y')$ of the fluorescent or of the bright field picture. Moments up to the 5 <sup>th</sup> order were calculated.
Eccentricity (Ecc) and I-Eccentricity (I-Ecc).	$(I-)Ecc = \sqrt{(I)\mu'_{20}/(I)\mu'_{02}}$ , were (I) $\mu'$ are the invariant second order normalised moments (Gonzales and Woods 2002).

*Principal Component Analysis (PCA) of FC-, QFM-, and QMM-data:* Fluorescent and morphologic parameters were determined for di- and triploids erythrocyte populations as described above.

The parameters  $P_i$  of the di- and triploid population were centred and normalised by

$P'_i = ((P_i - \bar{P}) / sd(P))$ . Eigen Values ( $\lambda$ ) and Eigen Vectors ( $V$ ) were calculated for each cell population (table 2). The probability that the members of each cell population are sorted correctly into the corresponding group (a and b) was quantified by discriminating capacities ( $C_D$ ,  $D = X/X, Y/X, Y/Z$  for 1/2/3 dimensions, figure 2):

$$C_D = \sum_D (d_a)_D \cdot \left( \frac{(d_a)_D}{(d_a)_D + (d_b)_D} \right) \quad [4]$$

signal to noise ratios ( $SNR_{DI/TRI}$ ) of the most discriminant parameters of FC, QFM, and QMM:

$$SNR_{DI/TRI} = \frac{|\bar{x}_{DI} - \bar{x}_{TRI}|}{sd(x_{DI/TRI})} \quad [5]$$

For the classification of unknown cell populations, parameters were transformed into the coordinate system of each calibration set ( $a$  and  $b$ ). Then, the Mahalanobis Distance (MD) (Taguchi and Jugulum, 2002) was calculated for each cell in respect to each coordinate system and the minimum distance criteria  $\min(MD_a, MD_b)$  was used to classify the cell into group  $a$  or  $b$ .

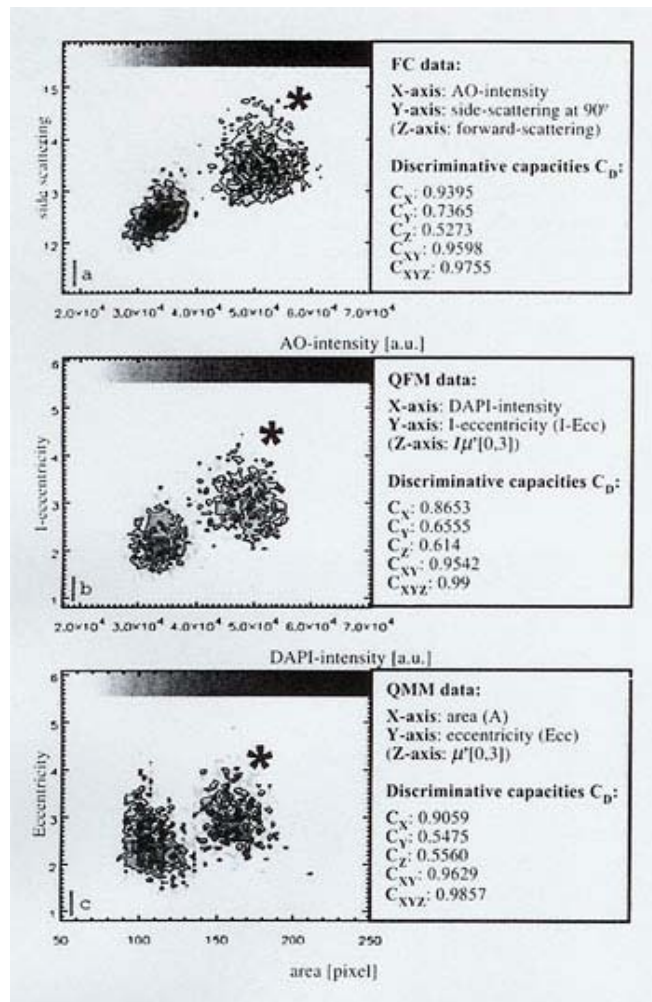
$$MD_{a,b} = \sum_i (P_{(a,b)i}) \quad [6]$$

## RESULTS

Segmentation of erythrocyte nuclei of *O. mykiss* from digital images was achieved in the bright field mode and in the fluorescent mode of the microscopic setup. [Figure 1](#) shows the basic steps of a representative segmentation of GEMSA-stained triploid nuclei which are also valid for the segmentation of diploid nuclei (results not shown). First, a threshold-interval is selected interactively in the intensity-histogram of the digital image ([figure 1a](#) and [1b](#), inset). The segmented regions of interest (white pixels in [figure 1b](#)) include erythrocyte nuclei, but also cellular debris and membrane residues. Two morphologic filters based on object size and circularity (shape sensitive parameter  $P^2/A$ , see [table 1](#)) separate the nuclei from the undesired structures ([figure 1c](#)). The insets of figure 1c show that cellular debris and membrane residues are smaller and less circular (high  $P^2/A$ -values) than the nuclei.

Morphological and intensity dependent parameters of segmented nuclei were calculated and the data were analysed by multi-parameter techniques ([table 1](#) and [2](#), [figure 2](#)). As [figure 2](#) shows, di- and tri-ploid (\*) erythrocyte populations were separated by FC ([figure 2a](#)), QFM ([figure 2b](#)), and QMM ([figure 2c](#)). The results are plotted in bivariate histograms with area normalised density distributions. All three methods clearly distinguish between the diploid and the triploid erythrocyte population. The corresponding  $C_D$ -values were calculated for 1-, 2-, and 3-dimensions and plotted next to each histogram. The  $C_X$ -values are highest for the FC-data (AO-intensity), followed by the QMM-data (area), and the QFM-data (DAPI-intensity). For QFM, staining with DAPI, PI, or AO yielded comparable results (not shown). For the 2- and 3-dimensional analysis, the  $C_D$ -values are almost identical. The calculation of the SNR with respect to the most discriminant parameters yielded:  $SNR_{DI/TRI} = 4.5/3.1$  (AO-intensity for FC),  $SNR_{DI/TRI} = 2.5/2.1$  (DAPI-intensity for QFM), and  $SNR_{DI/TRI} = 4.0/2.0$  (area for QMM).





**Figure 2.** Comparison of FC [a], QFM [b], and QMM [c] based multi parameter analysis of di- and triploid erythrocyte nuclei of *O. mykiss*. Bivariate histograms show area normalised density distributions with contour lines. Triploid nuclei are marked by \*. The maximal density in each histogram is set to the maximal grey level of the colour bars at the top of each plot. Densities were calculated in 86 intervals for each dimension. At the right side of each plot, the  $C_D$ -values are listed for the most discriminant parameters in X, Y, and Z (Z-axis not shown in the plots).

Comparación entre los parámetros obtenidos por FC [a], QFM [b] y QMM [c]. Los histogramas bivariados contienen líneas de contorno para las poblaciones de eritrocitos di- y triploides (\*). La densidad máxima en cada histograma fue escalada con el valor máximo de gris según la escala en el margen superior de cada gráfico. Las densidades fueron determinadas en 86 intervalos en cada dimensión. Las capacidades de discriminación ( $C_D$ ) para los ejes X, Y, y Z (no graficado) se indican a la derecha.

Additional insight in the interdependence between the parameters is obtained by multi parameter analysis of the FC-, QFM-, and QMM-data (table 2). The correlation coefficients ( $r$ ) between the X-, Y-, and Z-axis parameters (figure 2) are plotted in the first row of table 2. The correlation coefficient for the X/Y-axis of the FC-data is higher than the correlation coefficients of the QFM- and the QMM-data. This explains why the  $C_{XY}$ -values for the QFM- and the QMM-data finally vary little from the  $C_{XY}$ -value of the FC-data, although the  $C_X$ - and the  $C_Y$ -values of the FC-data are higher than the corresponding values for the QFM- and the QMM-data. The second column of table 2 shows the correlation coefficients calculated between fluorescence intensity-dependent parameters (I and I-Ecc) and pure morphological parameters (A, P, D,  $m'_{22}$ , and Ecc) of the QFM-data. Since  $r$ -values are close to one, fluorescence dependent parameters can be substituted by pure morphological parameters; the latter can be obtained by ordinary light microscopy and QMM. The third column of table 2 shows the

reduce the dimensions of the matrix for PCA-analysis to 13 x 13 (grouped parameters contain redundant information). PCA can be applied to the QMM-data since the extracted parameters yield a gauss-like distribution. Due to the different nuclear morphology, parameter groups formed by the diploid cell population differ in part from the groups formed by the triploid set. In consequence, the compositions of the first three principal axes which result from the PCA ([table 2](#), bottom) are similar but not identical for the di- and the tri-ploid erythrocyte populations (only parameter groups with a participation > 10% were plotted). For both erythrocyte populations, the calculated Eigen Values ( $\lambda$ ) for the first principal axis ( $\lambda_1$ ) are more than twice as high as those for the second principal axis ( $\lambda_2$ ). The ratio of the standard deviation ( $sd = \sqrt{\lambda}$ ) of the diploid nuclei in relation to the first two principal axis  $\sqrt{\lambda_1} / \sqrt{\lambda_2} = 1.47$  defines the relative importance of the major in respect to the minor principal axis for the classification of sample erythrocytes. This outlines the advantage of the Mahalanobis Distance over the simple Euclidean Distance: The weights compensate for the nonhomogeneous distributions of the cellular parameters in relation to each principal component. Accordingly, the Mahalanobis Distance represents a reliable measure for the classification of unknown objects into well-known populations.

**Table 2.** Principal component analysis of FC-, QFM-, and QMM-data. The first row contains the correlation coefficients ( $r$ ) between the X-, Y-, and Z-axis parameters used to calculate the discriminating capacities  $C_D$  in figure 2. The second row contains  $r$  calculated for the intensity dependent and pure morphological parameters of the QFM-data. The third row contains parameter groups (A-M) formed by parameters with  $r > 0.95$  of the QMM-data. The last row contains the participation of these parameter groups (A-M) in the composition of the first three principal axis (Eigen Vectors). For the first three principal axes, the Eigen Values ( $\lambda_i$ ) represent the importance of each axes for the calculation of the Mahalanobis Distance.

Análisis de componentes principales para datos de FC, QFM y QMM. La primera fila contiene los coeficientes de correlación ( $r$ ) entre los ejes X, Y y Z, utilizados en cálculo de la capacidad discriminatoria  $C_D$  en la figura 2. La segunda fila contiene  $r$  calculado a partir de parámetros dependientes de la intensidad y parámetros puramente morfológicos obtenidos por QFM. La tercera fila agrupa parámetros obtenidos por QMM que presentan un  $r > 0.95$  (A-M). La última fila contiene el porcentaje de participación de estos grupos (A-M) en la composición de los primeros tres ejes principales (Auto Vectores). Para los primeros tres ejes principales los Auto Valores ( $\lambda_i$ ) representan la importancia de cada eje en el cálculo del criterio de la mínima distancia según Mahalanobis.

	FC	QFM	QMM
Correlation coefficient ( $r$ ) between X-/Y-/Z-axis parameters, figure 2	X/Y: $r = 0.5837$ X/Z: $r = 0.0723$ Y/Z: $r = 0.3594$	X/Y: $r = 0.3637$ X/Z: $r = 0.1944$ Y/Z: $r = 0.1749$	X/Y: $r = 0.0545$ X/Z: $r = 0.3443$ Y/Z: $r = 0.1124$
	QFM		
	Diploid cells	Triploid cells	
Correlation coefficient ( $r$ ) between intensity dependent and morphological parameters	I/A: $r = 0.998$ I/D: $r = 0.985$ I/P: $r = 0.963$ I-Ecc/Ecc: $r = 0.997$	I/A: $r = 0.998$ I/D: $r = 0.969$ I/ $\mu'_{22}$ : $r = 0.964$ I-Ecc/Ecc: $r = 0.998$	
	QMM		
	Diploid cells	Triploid cells	
Groups formed by the morphologic parameters with correlation coefficients $r > 0.95$	A: [ $\mu'_{22}$ ], [ $\mu'_{41}$ ] B: $E_C$ , $E_{C'}$ , $2^{nd}_d$ , $2^{nd}_{d'}$ C: [ $\mu'_{01}$ ] D: $\underline{D}$ E: A F: D G: [ $\mu'_{40}$ ] H: [ $\mu'_{01}$ ], [ $\mu'_{05}$ ] I: [ $\mu'_{33}$ ], [ $\mu'_{55}$ ] J: [ $\mu'_{30}$ ], [ $\mu'_{50}$ ] K: Ecc L: $P^2/A$ M: P	A: A, D, [ $\mu'_{22}$ ], [ $\mu'_{44}$ ] B: $E_C$ , $E_{C'}$ C: [ $\mu'_{01}$ ] D: $\underline{D}$ E: $2^{nd}_d$ , $2^{nd}_{d'}$ F: [ $\mu'_{33}$ ] G: [ $\mu'_{40}$ ] H: [ $\mu'_{01}$ ], [ $\mu'_{05}$ ] I: [ $\mu'_{55}$ ] J: [ $\mu'_{30}$ ], [ $\mu'_{50}$ ] K: Ecc L: $P^2/A$ M: P	
Participation of the parameter groups A-M in the 1 <sup>st</sup> , 2 <sup>nd</sup> , and 3 <sup>rd</sup> principal axes together with the size of the Eigen Values ( $\lambda$ ) of each axis in relation to the sum of all $\lambda$ s	1 <sup>st</sup> axis, $\lambda_1 = 0.42$ : A: 16.8%, C: 11.9%, E: 16.3%, F: 15.2%, M: 16.4% 2 <sup>nd</sup> axis, $\lambda_2 = 0.2$ : G: 10.0%, H 11.1%, K: 28.4%, L: 25.9% 3 <sup>rd</sup> axis, $\lambda_3 = 0.1$ : I: 50.3%, J: 48.9%	1 <sup>st</sup> axis, $\lambda_1 = 0.38$ : A: 17.1%, C: 15.5%, G: 12.8%, M: 18.0% 2 <sup>nd</sup> axis, $\lambda_2 = 0.16$ : F: 42.9%, I: 43.1% 3 <sup>rd</sup> axis, $\lambda_3 = 0.14$ : B: 11.5%, D: 14.0%, E: 15.8% K: 26.7%, L: 12.0%	

## DISCUSSION

In 1991 Kamentsky and Kamentsky reported that quantitative cytometric data obtained with fluorescence microscopy is comparable to data obtained with FC. Many different applications have emerged since then which take advantage of quantitative fluorescence microscopy or quantitative morphological microscopy, QFM or QMM (e.g. Fanani *et al* 2002, Härtel *et al* 2003/2005, Alvarez *et al* 2004/2005). As we report here, the relative DNA content of erythrocytes of *O. mykiss* can be quantified with FC (AO-staining) or QFM (AO-, DAPI-, or PI-staining). The discriminating capacities  $C_D$  indicate that the relative DNA content in erythrocytes can be determined by QFM with a precision close to the FC-data (the reference method in quantitative cytometry). Unfortunately, determination of morphological parameters with FC is not reliable; side- and the forward-scattering of the laser beam resemble surface

or QMM-data, and we observe that the  $C_{XY}$ - and  $C_{XYZ}$ -values for QFM and QMM catch up to or are even greater than the FC-data. In consequence, the strength of the image processed microscopic methods QFM and QMM lies in the precise determination of more than one parameter coupled to the fluorescence (QFM) or to the morphology (QMM) of objects.

To our surprise, the integral fluorescence intensities of the di- and triploid erythrocyte nuclei correlate almost perfectly with basic morphologic properties ( $r > 0.99$  for I and A). The correlation between the relative DNA content and the nuclear size suggests a homogeneous packing of the chromatin inside the nuclear membrane. In consequence, the relative DNA content in erythrocytes can be estimated indirectly by the size of the nuclei. This conclusion reveals why cell size and morphology have been used as discriminant parameters for the identification of di- and triploid salmon populations (Phillips *et al* 1986, Wlasow *et al* 2004). In addition to the correlation between relative DNA content and the nuclear size, further intensity coupled parameters can be substituted by their morphologic counterparts ( $r > 0.99$  for I-Ecc and Ecc). Since morphologic parameters are calculated directly from the binary mask of the objects, fluorescence staining of the nuclei becomes unnecessary and can be substituted by simple, inexpensive, and non hazardous contrast staining procedures like GIEMSA in combination with bright field microscopy. For the morphological discrimination between erythrocyte populations with QMM, size, eccentricity (a measure for circularity), and invariant moments were shown to be the most accurate parameters (table 2, figure 2). This result is supported by the visual impression: diploid nuclei are small and circular in comparison to the curved, bean-like triploid nuclei; see Small and Benfey (1987) for diploid erythrocytes and figure 1 or Wlasow *et al* (2004) for triploid samples.

Without PCA, visual multi-parameter analysis becomes a difficult task when more than two parameters are considered. For  $n$  parameters,  $n(n-1)/2$  two-dimensional histograms have to be monitored in order to find an optimal representation. With increasing  $n$ , the task becomes time-consuming if not impossible. As shown, QMM in combination with PCA performs excellent feature extraction for di- and triploid erythrocyte populations (table 2, bottom). PCA bundles redundant information and reduces the dimension of the parameter space to a minimum. Besides PCA, alternative methods like principal component regression, partial least square regression, and artificial neuronal networks have been tested by Davey *et al* (1999) for FC classification of different micro-organism with up to 7 parameters. The authors reported that artificial neuronal networks provided the best discriminating capacities in most experiments. For QMM, we preferred PCA in combination with the Mahalanobis Distance, because it presents a very clear classification criterion.

In conclusion, QMM can quantify triploidization rates in rainbow trout in a reliable, fast, and innovative way. The results are comparable or even superior to the existing reference method, FC. Since di- and tri-ploid rainbow trout do not show significant variations of erythrocyte density in the blood (data not shown), aliquots of blood samples can be mixed into a single sample. This way, the time and cost-intensive analysis of many individual blood samples can be reduced to a single step. The developed method opens a perspective for the classification of microscopic objects with many possible applications.

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