

Sperm swimming speed and morphology differ slightly among the three genetic morphs in the ruff sandpiper (*Calidris pugnax*), but show now clear polymorphism

by [Martin Bulla](#), Clemens Küpper, David B Lank, Jana Albrechtová, Jasmine L Loveland, Katrin Martin, Kim Teltcher, Margherita Cragolini, Michael Lierz, Tomáš Albrecht, Wolfgang Forstmeier & Bart Kempenaers

Methods

S0 – Deviations from the *a priori* protocols

Sperm competition

We initially based our predictions about between-morph differences in sperm traits on the idea that the intensity of sperm competition might be stronger in Faeders and might thus have selected for more competitive sperm in this morph. However, we realized, that in ruffs the likelihood that beneficial mutations occur is several orders of magnitude lower in Faeders than in Independents. We now discuss this in the manuscript and also refrain from predictions about better sperm as what characterizes high quality sperm in birds is debated (see references in the main text).

Vas deference samples

We have not included the morphology data based on the vas deference samples, because (a) we did not have corresponding sperm velocity measurements, (b) we were unsure whether those sperm were fully developed and (c) we used a different method ([ImageJ](#)) to measure sperm morphology than the semi-automated Sperm sizer method that was developed later and that we used for the sperm collected by abdominal massage, i.e. the measures are not directly comparable.

Velocity measurements

We aimed at using curvilinear velocity as a measure of sperm speed, but report all three velocities provided by the sperm tracking software, for reasons described in section S2 below. We suggest that straight-line velocity is a better velocity measure in ruffs.

Coefficient of variation

Our main goal was to investigate how the within-male coefficient of variation in sperm length differs between the morphs. We did not use the between-male coefficient of variation per morph because of the unbalanced number of males per morph within our sample.

S1 - Housing

Individuals were housed in groups of varying size and different morph and sex composition (Table S0), in two outdoor aviaries (123 m² and 119 m²) and in a complex of semi-outdoor aviaries divided into a large central space (117 m²) and 24 smaller, adjacent aviaries (8.8 m² each). One aviary contained only males and in three aviaries male access to females was limited to 3.5h per day. Note that males in the male-only aviaries also produced sperm, and we observed homosexual matings in this setting. The semi-outdoor aviaries have a solid but transparent roof, and wire-mesh on at least one side, so that the birds experienced natural light and temperature cycles. All aviaries had natural grass, a few wooden logs, small elevated areas suitable for lekking, and at least one heatable water body (~1 m²). Aviaries were cleaned daily and the birds were provided with a mixture of food pellets for waders, dried shrimp and live mealworms ([Meghlys](#); www.meghlys.com) in dishes or spread over the grass.

Table S0 | Distribution of male morphs and their access to females across aviaries

Location	Access to ♀	Aviary	N ♀	N ♂	Independents	Satellites	Faeders
A	always	1	15	3	3		
	always	2	8	3	3		
	always	3	8	3	1	2	
	always	4	15	3	3		
B	3.5h/day	1	10	7	5	2	
	3.5h/day	2	10	10	8	2	
	none	3	8	8	2	5	1
	always	4	19	5	5		
	always	5	19	5			5
	always	6	19	5		5	
C	always	1-3	21	40	29	9	2

S2 - Velocities

During recording sessions, we observed that ruff sperm swim slower, i.e. cover shorter distances per unit of time, than passerine sperm ([video examples](#)). However, curvilinear velocity, our intended measure of velocity and a measure of sperm swimming speed typically used in studies on passerines (Laskemoen *et al.* 2010, Cramer *et al.* 2016, Opatová *et al.* 2016, Tomasek *et al.* 2017, Støstad *et al.* 2018, Schmoll *et al.* 2020), tracks the sperm's sideways "vibration" movements, and suggests that ruff sperm moves faster than passerine sperm (Fig. [S1](#)). Thus, in ruffs, the curvilinear velocity method likely does not reflect the actual velocity in terms of "distance covered per unit of time" and hence might not be the most appropriate velocity measure. Straight-line velocity seems more appropriate and reflects our observation that ruff sperm moves forward more slowly than zebra finch sperm. Straight-line velocity has also been linked to fertilization success in avian and non-avian taxa, and has been used in previous studies instead of or along with curvilinear velocity (reviewed in Pizzari *et al.* 2004, Denk *et al.* 2005, Helfenstein *et al.* 2009). Consequently, we report all three velocity measures provided by the sperm tracking software (curvilinear, straight-line, and average-path velocity). The three velocity measures are correlated ($r = 0.39 - 0.91$, Fig. [S2](#)). The within-male seasonal repeatability of sperm velocity, i.e. the percentage of the variation attributed to variation among males, was 24% for straight-line velocity, 35% for average-path velocity and 47% for curvilinear velocity (based on May and June velocity estimates of the same individuals; Fig. [S3](#) and [S4](#), Table [S1](#)).

S3 - Relationships between sperm components

In ruffs, sperm head length constitutes on average 23% of the total sperm length (range: 18 – 34%, $N = 920$ sperm from 92 males), the midpiece 17% (15 – 22%) and the tail 60% (45 – 65%). The head length reflects the length of the nucleus ($r = 0.97$), which makes up 82 – 96% of the head, while flagellum and total sperm length mainly reflect tail length ($r = 0.95$ and 0.89 respectively; Fig. [S5](#)). Head length did not correlate with midpiece length ($r = 0.00$) or tail length ($r = -0.02$), and midpiece length and tail length correlated weakly ($r = 0.27$; Fig. [S5](#)). The within-male within sperm-sample repeatability of sperm morphology measures varied from 25% to 60%, with acrosome length, the smallest part of the sperm, being the least repeatable (Fig. [S3](#), Table [S1](#)). We did not measure between-sample repeatability in sperm components as avian sperm is highly heritable (Birkhead *et al.* 2005).

S4 – Inbreeding and Relatedness

Because of our captive population, we expect higher levels of inbreeding compared to males from the wild. Previous studies on birds and mammals have shown that inbred males have lower sperm velocity and a higher proportion of abnormal sperm than outbred males (Gomendio *et al.* 2000, Ala-Honkola *et al.* 2013, Heber *et al.* 2013, Opatová *et al.* 2016). However, there is no evidence that the morphology of normal-looking sperm (e.g., length, coefficient of variation) differs between inbred and outbred males (Mehlis *et al.* 2012, Ala-Honkola *et al.* 2013, Opatová *et al.* 2016). Based on these studies, we assumed that our sperm velocity measurements may be on average somewhat lower than those of free-living ruffs, whereas the morphological measurements likely reflect the variation in sperm morphology observed in the wild. Because the morphs interbreed in our aviary, inbreeding levels between morphs should be similar. To assess whether inbreeding and relatedness between individuals influenced our results, we estimated these traits based on genotypes of all males at 21 polymorphic microsatellite markers (Giraldo-Deck *et al.* 2022).

We quantified inbreeding as homozygosity-by-locus (Aparicio *et al.* 2006) using the 'GENHET' R-function version 3.1 (Coulon 2010) and calculated Pearson's correlation coefficients between sperm traits and homozygosity. Correlations were weak (mean $r = -0.09$, range: -0.26 to 0.07 ; Fig. [S7](#)), and particularly so for measures of velocity ($r_{\text{curvilinear}} = -0.04$, $r_{\text{straight line}} = 0.07$, $r_{\text{average path}} = 0.05$), so we did not control the subsequent models for inbreeding.

To investigate whether the main results were confounded by relatedness between some of the individuals, we tested for a relatedness signal in the model residuals. We specified the model residuals as a new response variable in an intercept-only Bayesian linear regression fitted in STAN (Stan-Development-Team 2022) using the 'brm' function from the 'brms' R-package (Bürkner 2017, Bürkner 2018, Bürkner 2021) with the male relatedness matrix as a random effect. We constructed the relatedness matrix from the genotypes at 21 microsatellite loci using the 'coancestry' function from the 'related' R-package (Pew *et al.* 2015) and its "lynchli estimator", which correlated best with the expected values (although by a small margin). As the software sometimes estimates negative values for zero relatedness, negative relatedness values were assigned as zero. To make the relatedness matrix positive definite we added 0.1 to its diagonal. We used the default 'brm' priors, i.e. a flat prior for the intercept and a Student's t distribution for the standard deviation (Bürkner 2017, Bürkner 2018, Bürkner 2021). To decrease the possibility of divergent transitions threatening the validity of posterior samples, the target average proposal acceptance probability was increased to 0.99 or to 0.999 for the coefficients of variation (Bürkner 2017, Bürkner 2018, Bürkner 2021). Four Markov chains ran for 50,000 iterations each. For each chain, we discarded the first 25,000 iterations

and sampled every 20th iteration, which resulted in a total of 5,000 samples (4 x 1,250) of model parameters. We assessed the independence of samples in the Markov chain using graphic diagnostics and the convergence using the Gelman-Rubin diagnostics, which was 1 for all parameters, indicating model convergence (Brooks & Gelman 1998).

The relatedness matrix explained little variation in the residuals of models on swimming speed (10% for curvilinear, 5% for straight-line and 8% for average-path velocity) and close to zero variation in the residuals of all models on morphology (0.1%; Table S3). The alternative models using all velocity recordings or using individual sperm measurements gave similar results (Table S3). Importantly, the intercept only models without the relatedness matrix fitted the residuals better than the intercept only models with the relatedness matrix as a random effect; the estimated Bayes factor in favour of the model without the relatedness matrix ranged from seven to infinity (Lee & Wagenmakers 2014), and the posterior probability from 0.88 to 1 (mean = 0.97; Table S3). Thus, in the main text we report results from models without control for relatedness. In contrast, the relatedness matrix explained variable amounts of variation in the residuals from models on coefficients of variation in sperm morphology (7-77%, Table S3). As including the relatedness matrix did not change the conclusions (Fig. S8) and for consistency, we did not control for relatedness in the models described in the main text.

Figures

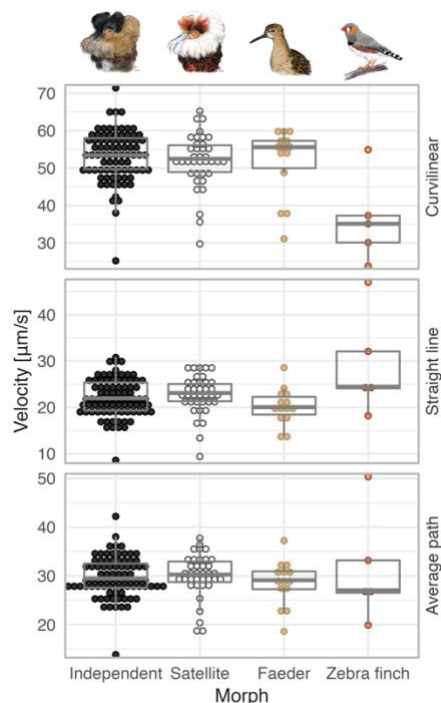


Figure S1 | Comparison of sperm swimming speed between ruff morphs and zebra finch. Dots represent velocity values for sperm of 4 ruff and 5 zebra finch males recorded in May, 46 ruff males recorded in June, and two values for 42 ruff males recorded in May and June. Boxplots depict median (horizontal line inside the box), the 25th and 75th percentiles (box) and the 25th and 75th percentiles ± 1.5 times the interquartile range or the minimum/maximum value, whichever is smaller (bars). Five zebra finch males from a population at the Max Planck Institute for Biological Intelligence in Seewiesen were sampled in June along with the ruffs to ensure that the ruff motilities and velocities are not an artefact of our sampling method. The zebra finch sperm moved normally (see [example](#)), with velocity values well within the range of measurements from our laboratory (Opatová *et al.* 2016, Knief *et al.* 2017). Note that the zebra finches were sampled outside a breeding phase (no nest boxes available). Created with 'ggplot' function and dots stacked using 'geom_dotplot' function, both from the 'ggplo2' R-package (Wickham 2016). Illustrations by Yifan Pei under [Creative Commons Attribution \(CC BY 4.0\)](#).

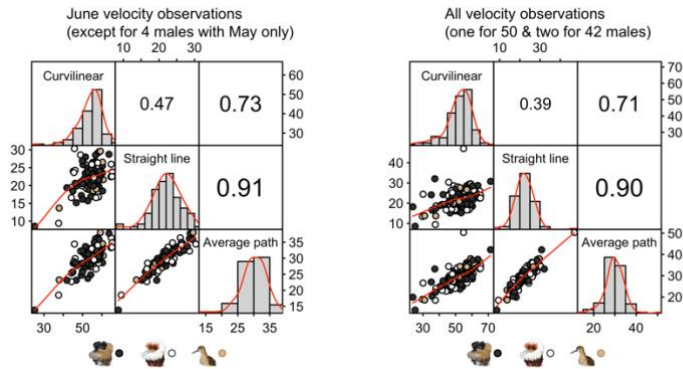


Figure S2 | Pairwise correlations among sperm velocity measures of ruffs. On the diagonal: histograms and density lines (red) for each variable. Above diagonal: Pearson's correlation coefficients with size highlighting the strength of the correlation. Below diagonal: bivariate scatterplots, with each dot representing June value per male (left), except for four males with May only values ($N = 92$), or all velocity observations (right; $N = 134$), dot color highlighting morph (black: Independents, white: Satellites, beige: Faeders) and red line representing loess-smoothed fit. Adapted from 'pairs.panels' function from 'psych' R-package (Revelle 2022). Ruff morph illustrations by Yifan Pei under [Creative Commons Attribution \(CC BY 4.0\)](https://creativecommons.org/licenses/by/4.0/).

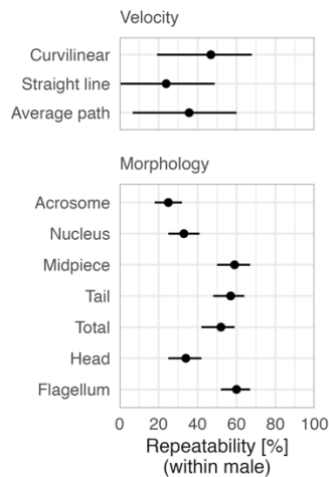


Figure S3 | Within-male repeatability of sperm traits in ruffs. Dots with bars represent repeatability estimates with 95%CI generated by the 'rpt' function from the 'rptR' R-package, without specifying fixed effects (Stoffel *et al.* 2017). For sperm velocity we used one measurement from May and one from June ($N = 42$ males). For sperm morphology, we used measurements of 10 sperm per male ($N = 92$ males). The last three morphological traits are composite traits (Total = Acrosome + Nucleus + Midpiece + Tail, Head = Acrosome + Nucleus, Flagellum = Midpiece + Tail). For precise estimates see Table [S1](#).

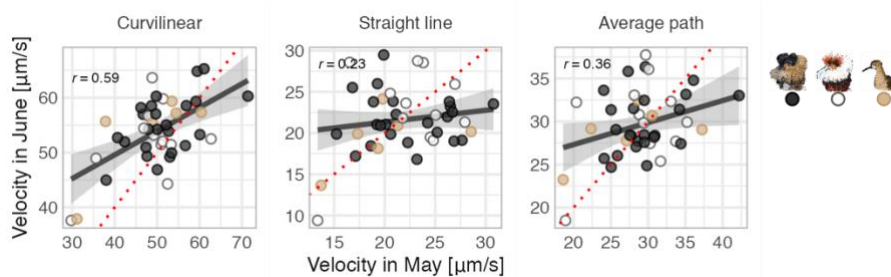


Figure S4 | Correlation between May and June sperm velocity of ruffs. Dots represent single males, dot color the morph (black: Independents, white: Satellites, beige: Faeders), lines with shaded areas linear model fits with 95%CI generated by 'stat_smooth' function in 'ggplot2' R-package (Wickham 2016) using robust regression specified by 'rlm' function from 'MASS' R-package (Venables & Ripley 2002). 'r' represents Pearson's correlation coefficient and dotted lines indicate equality, i.e. points above the line represent faster sperm in June, points below the line faster sperm in May. $N = 42$ males with velocity measured both in May and June (24 Independents, 12 Satellites, 6 Faeders).

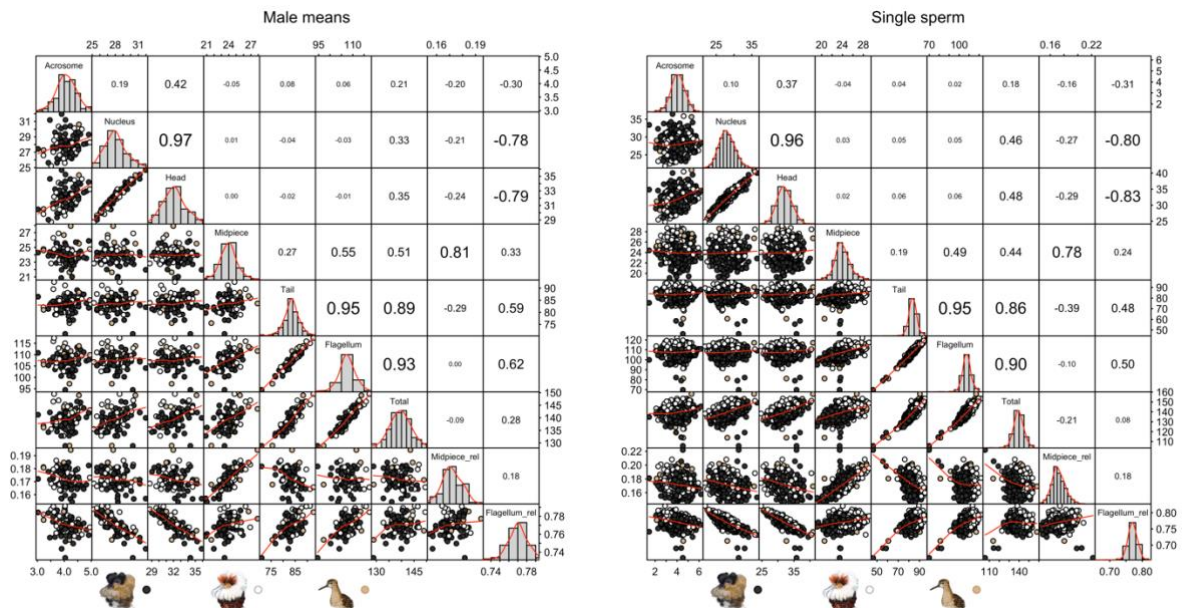


Figure S5 | Pairwise correlations among sperm morphological traits of ruffs. On the diagonal: histograms and density lines (red) for each variable. Above diagonal: Pearson's correlation coefficients with size highlighting the strength of the correlation. Below diagonal: bivariate scatterplots, with each dot representing average value per male (left; N = 92) or a single sperm value (right; N = 920), dot color highlighting morph (black: Independents, white: Satellites, beige: Faeders) and red line representing loess-smoothed fit. Adapted from 'pairs.panels' function from 'psych' R-package (Revelle 2022). Ruff morph illustrations by Yifan Pei under [Creative Commons Attribution \(CC BY 4.0\)](#).

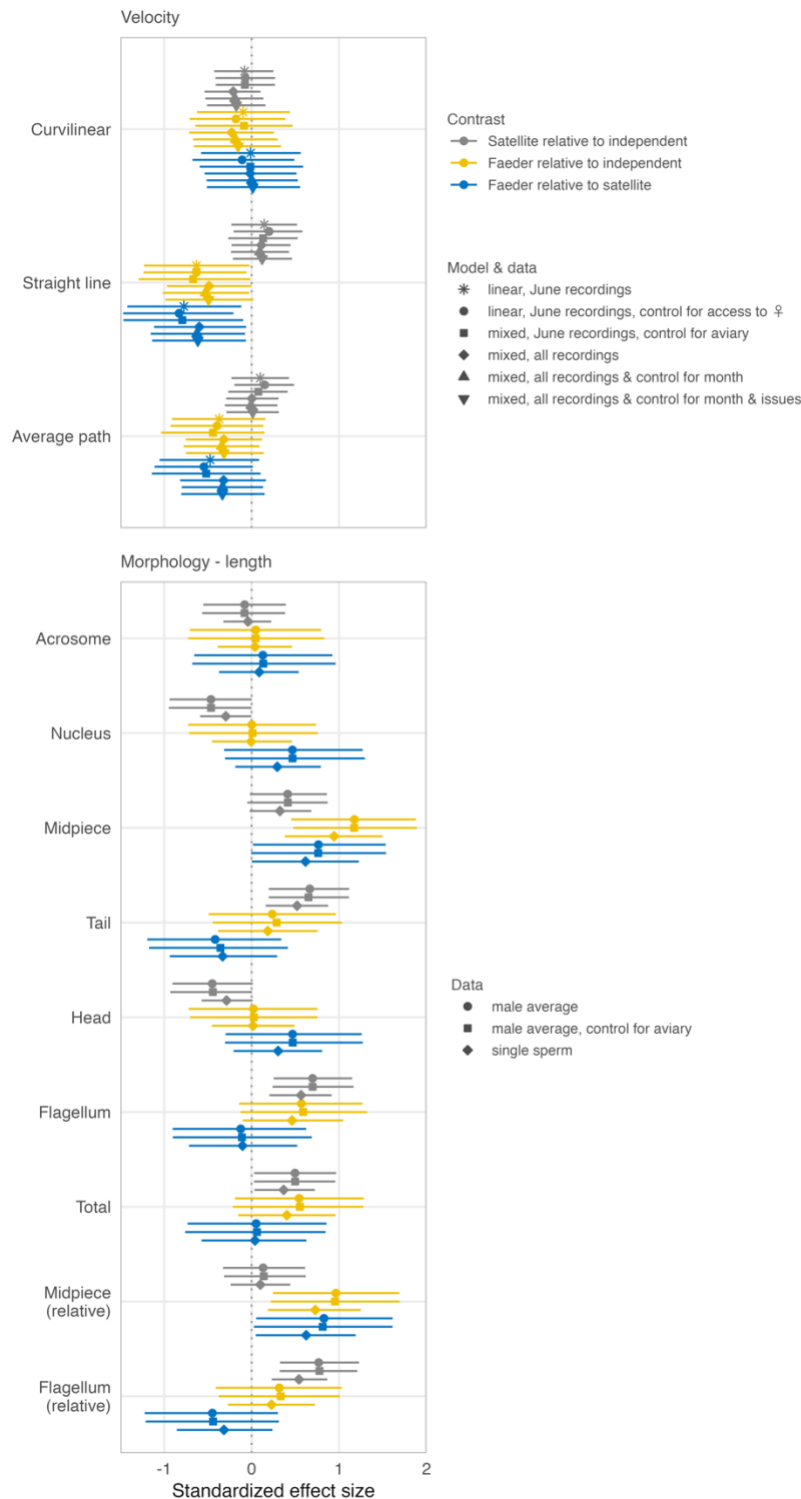


Figure S6 | Differences in sperm traits of ruff morphs according to model and data type. Shapes with bars represent estimated standardized effect sizes (medians) with their 95% CIs based on the joint posterior distribution of 5,000 simulated values generated from models by the 'sim' function from the 'arm' R-package (Gelman & Su 2021). Color highlights the between-morph differences – Satellite relative to Independent (grey), Faeder relative to Independent (yellow), Faeder relative to Satellite (blue) – and shape indicates type of model and data. For **velocity**, '*linear, June recordings*' indicates results based on a linear model fitted to the June velocity values of all males, except for four males for which only recordings from May were available; '*mixed, June recordings, control for aviary*' indicates results based on the same data but from a mixed-effect model that includes 'aviary' as a random intercept; '*mixed, all recordings*' indicates results based on a mixed-effect model fitted to all velocity values (including 42 males with a recording for both May and June) with male identity included as a random intercept. To control for 'access to ♀' (none, temporal, continuous), 'month' (May or June) and/or 'issues' (yes or no; indicating e.g. presence of faeces) in some models we included these terms as fixed effects. For **morphology**, '*male average*' indicates results based on linear models fitted to the average morphological value per male based on 10 sperm cells; '*male average, control for aviary*' indicates results based on the same data but from mixed-effect models containing aviary as a random intercept; '*single sperm*' indicates results based on mixed-effect models fitted to single sperm-cell measurements (10/male) and controlled for multiple sampling per male by including male identity as a random intercept. For both, velocity and morphology, the models from the main text Fig. 2 are listed first and in the above legends indicated as 'linear, June recordings' and 'male average'. Note that the main text estimates are similar to those from the alternative models.

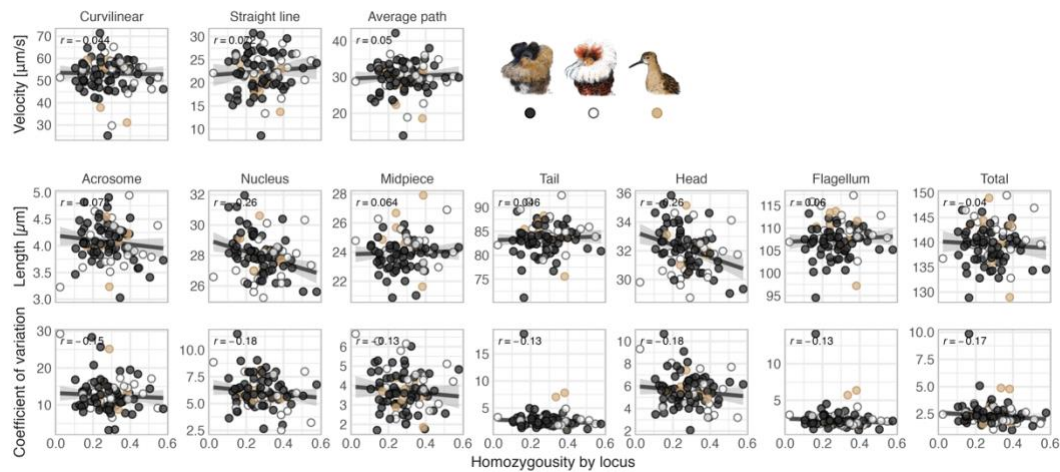


Figure S7 | Correlation between sperm traits and homozygosity by locus. Dots represent single male velocity values from June (except for four males with May only values) or male average lengths and coefficients of variation from 10 sperm cells (N = 92 males). Dot color indicates morph: black – Independent, white – Satellite, beige – Faeder. Lines with shaded area represent linear model fit with 95%CI generated by 'stat_smooth' function in 'ggplot2' R-package (Wickham 2016) using robust regression specified by 'rlm' function from 'MASS' R-package (Venables & Ripley 2002). 'r' represents Pearson's correlation coefficient. Ruff morph illustrations by Yifan Pei under [Creative Commons Attribution \(CC BY 4.0\)](https://creativecommons.org/licenses/by/4.0/).

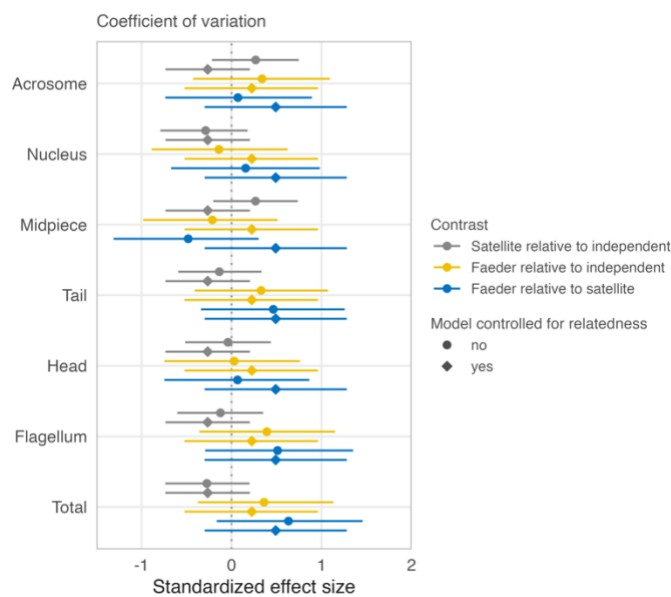


Figure S8 | Ruff morph differences in coefficients of variation of sperm traits with and without control for relatedness. Shapes with bars represent estimated standardized effect sizes with their 95%CI, color the between-morph differences – Satellite relative to Independent (grey), Faeder relative to Independent (yellow), Faeder relative to Satellite (blue) – and shape indicates whether model was controlled for relatedness (diamond) or not (dot). Note that the estimates reported in the main text (Fig. 2, here indicated by dots) are similar to those from models controlled for relatedness (diamonds). Estimates (medians) and 95%CI are based on the joint posterior distribution of 5,000 simulated values generated by the 'sim' function from the 'arm' R-package (Gelman & Su 2021) using the model outputs from Table S1 (dots) or generated by 'brm' function in 'brms' R-package (Bürkner 2017, Bürkner 2018, Bürkner 2021) with a vague (weakly informative) Gaussian priors centered on zero for the intercept and factor levels and half Cauchy priors for the standard deviations, including the error term - standard deviation of the residuals (diamonds). Using default 'brms' priors generated same results. For further details see Methods S1, for comparison of models with and without control for relatedness Table S3.

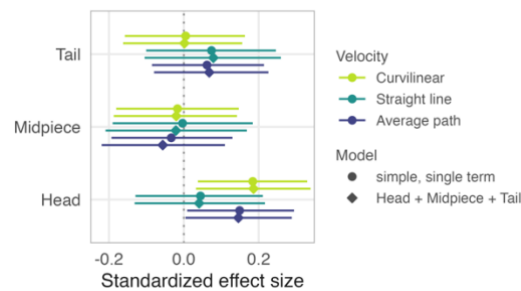


Figure S9 | Estimates from simple univariate models are similar to those from a model containing multiple traits. Shapes with bars represent estimated standardized effect sizes with their 95% CIs based on the joint posterior distribution of 5,000 simulated values generated from models by the 'sim' function from the 'arm' R-package (Gelman & Su 2021). Color indicates dependent variable, i.e. type of velocity (green: curvilinear, turquoise: straight line, purple: average path). Shape indicates source of estimate: from 'univariate' models with single morphological terms (head, midpiece or tail; dot) or a multivariate model containing all three terms (diamond). The models were controlled for number of tracked sperm (ln-transformed) and morph. The response (velocity) as well as the linear term/s (number of tracked sperm, head, midpiece and tail) were scaled (mean-centered and divided by standard deviation).

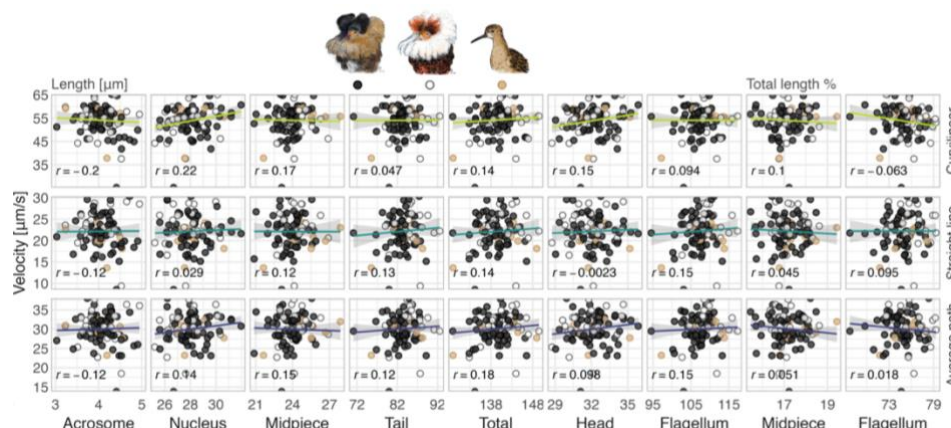


Figure S10 | Sperm swimming speed in relation to sperm morphology. Lines with shaded areas represent model predictions with their 95% CIs based on the joint posterior distribution of 5,000 predicted values generated from linear models, controlled for number of tracked sperm (ln-transformed) and morph (Table S5). 'r' represents Pearson's correlation coefficient (ignoring morph, hence r may differ from model estimates). Dots represent data points based on single June-values for velocity (with exception of four males with May-values only) and average trait lengths of 10 sperm cells per male, dot color highlights morph. Ruff-morph illustrations by Yifan Pei under [Creative Commons Attribution \(CC BY 4.0\)](https://creativecommons.org/licenses/by/4.0/).

Tables

Table S1 | Within male repeatability of ruff sperm traits

Trait	Specification	Estimate	95%CI
Velocity	Curvilinear	47%	19-68%
	Straight line	24%	0-49%
	Average path	36%	7-60%
Length	Acrosome	25%	18-32%
	Nucleus	33%	25-41%
	Midpiece	59%	50-67%
	Tail	57%	48-64%
	Head	34%	25-42%
	Flagellum	60%	53-67%
	Total	52%	42-59%

Repeatability estimates with 95%CI generated by 'rpt' function from 'rptR' R-package (Stoffel *et al.* 2017) from two - May and June - velocity estimates per male (N = 42 males) and 10 sperm morphology measurements per male (N = 92 males).

Table S2 | Within and between observer repeatability of ruff sperm measurements

Trait	Repeatability estimate (95%CI)	
	Within observer	Between observer
Acrosome	92% (85.2 - 95.6)	90% (80.8 - 94.5)
Nucleus	98% (97.1 - 99.1)	98% (95.6 - 98.7)
Midpiece	99% (98.5 - 99.6)	99% (98.7 - 99.6)
Tail	97% (94.3 - 98.5)	99% (97.3 - 99.2)
Total	100% (99.4 - 99.8)	99% (98.1 - 99.4)
Head	98% (95.6 - 98.8)	97% (94.3 - 98.4)
Flagellum	99% (97.5 - 99.3)	98% (96.4 - 98.9)

Repeatability estimates with 95%CI generated by the 'rpt' function from 'rptR' R-package (Stoffel *et al.* 2017) from 40 sperm measured twice by the same observer (within observer) or by two different observers (between observer). Note that the smallest part - acrosome - has the lowest repeatability, but the repeatability of head, the composite measure of acrosome and nucleus, is high.

Table S3 | Percentage of variation in ‘residuals from original models’ explained by relatedness and comparison of models without and with control for relatedness.

Data	Trait	Specification	Variance explained by relatedness	95% CI			Bays factor	Probability of model without relatedness matrix
June	Velocity	Curvilinear	10.1%	0.0%	44.7%	7	0.875	
		Straight line	5.1%	0.0%	34.7%	10	0.913	
		Average path	8.0%	0.0%	38.8%	8	0.886	
Averages	Length	Acrosome	0.1%	0.0%	0.7%	Inf	1	
		Nucleus	0.1%	0.0%	0.7%	Inf	1	
		Midpiece	0.1%	0.0%	0.6%	Inf	1	
		Tail	0.1%	0.0%	0.6%	Inf	1	
		Total	0.1%	0.0%	0.6%	Inf	1	
		Head	0.1%	0.0%	0.7%	Inf	1	
		Flagellum	0.1%	0.0%	0.6%	Inf	1	
		Male value	Coefficient of variation	Acrosome	33.9%	0.3%	77.1%	1
Nucleus	16.8%			0.1%	58.1%	3	0.767	
Midpiece	6.6%			0.0%	43.7%	7	0.874	
Tail	76.8%			24.0%	94.3%	0	0.1	
Total	47.8%			0.5%	85.6%	1	0.528	
Head	20.3%			0.1%	60.0%	2	0.724	
Flagellum	69.9%			3.4%	94.9%	0	0.278	
All	Velocity			Curvilinear	1.8%	0.0%	15.6%	22
		Straight line	1.9%	0.0%	16.9%	15	0.937	
		Average path	2.4%	0.0%	17.4%	16	0.948	
All	Length	Acrosome	0.1%	0.0%	0.7%	84	0.989	
		Nucleus	0.1%	0.0%	0.7%	94	0.99	
		Midpiece	0.1%	0.0%	0.6%	132	0.992	
		Tail	0.1%	0.0%	0.6%	118	0.991	
		Total	0.1%	0.0%	0.6%	113	0.992	
		Head	0.1%	0.0%	0.7%	90	0.989	
		Flagellum	0.1%	0.0%	0.6%	143	0.993	

Variance explained by relatedness with **95%CI** represents percentage of variance explained by relatedness matrix in an intercept only model fitted to residuals of the original models (Fig. 2, Table 1) in STAN (Stan-Development-Team 2022) using ‘bm’ function from ‘brms’ R-package (Bürkner 2017, Bürkner 2018, Bürkner 2021) with male relatedness matrix and male identification (in case of residuals from models on single values) as random effect. **Bayes factor** in favor of model without relatedness matrix and **probability of model without relatedness matrix** in comparison to a model with relatedness matrix. The Gelman-Rubin diagnostics was 1 for all models, indicating model convergence (Brooks & Gelman 1998). Note that for all cases, but coefficient of variation in Tail and Flagellum, the model without relatedness matrix fits residuals better than a model with relatedness matrix, which justifies our use of simple original models, not controlled for relatedness (Fig. 2). Importantly, controlling the original models on coefficient of variation for relatedness generated similar results (Fig S8).

Table S4 | AICc comparison of simple and quadratic effects of morphological traits on sperm velocity

Velocity	Trait	AICc of the model		Δ AICc	Akaike weight	Evidence ration
		Simple	Quadratic			
Curvilinear	Acrosome	208.44	210.59	2.15	0.03	32.33
	Nucleus	200.41	202.71	2.30	0.03	32.33
	Midpiece	209.02	209.89	0.87	0.06	15.67
	Tail	209.07	211.28	2.21	0.03	32.33
	Total	208.23	209.84	1.61	0.04	24
	Head	202.81	205.07	2.26	0.03	32.33
	Flagellum	209.07	211.33	2.26	0.03	32.33
	Midpiece relative	208.41	210.50	2.09	0.03	32.33
	Flagellum relative	204.84	206.93	2.09	0.03	32.33
	Acrosome	229.05	230.93	1.88	0.04	24
Straight line	Nucleus	228.75	231.05	2.30	0.03	32.33
	Midpiece	229.06	231.05	1.99	0.04	24
	Tail	228.30	230.55	2.25	0.03	32.33
	Total	228.25	228.88	0.63	0.07	13.29
	Head	228.77	231.05	2.28	0.03	32.33
	Flagellum	228.47	230.33	1.86	0.04	24
	Midpiece relative	228.67	230.60	1.93	0.04	24
	Flagellum relative	229.05	231.29	2.24	0.03	32.33
	Acrosome	202.34	204.64	2.30	0.03	32.33
	Nucleus	198.15	200.49	2.34	0.03	32.33
Average path	Midpiece	202.38	204.40	2.02	0.04	24
	Tail	201.80	204.14	2.34	0.03	32.33
	Total	200.74	201.93	1.19	0.05	19
	Head	198.34	200.68	2.34	0.03	32.33
	Flagellum	202.14	204.24	2.10	0.03	32.33
	Midpiece relative	200.68	201.97	1.29	0.05	19
	Flagellum relative	200.90	203.24	2.34	0.03	32.33

Each model was controlled for number of tracked sperm (ln-transformed) and morph. The response (velocity), number of tracked sperm and linear morphological term (in case of simple models) were scaled (mean-centered and divided by standard deviation). **Simple** represents AICc value for a model with a linear morphological term (scaled) and **Quadratic** AICc value for a model with a linear and quadratic morphological term (second polynomial). Δ **AICc** - the difference in AICc between the quadratic and simple model (i.e. positive values indicate a poorer fit of the quadratic model). **Akaike weight** - the weight of evidence that a quadratic model is the best approximating model, i.e. probability of the quadratic model. Note that the probability of the Simple model (with linear term only) is $1 - w_i$ and hence 0.93-97. **Evidence ration** - the model weight of the simple model relative to the quadratic model, i.e. how many times is the simple model more likely than the quadratic model. All parameters confirm superiority of the simple models.

Table S5 | Ruff sperm velocity in relation to sperm morphology.

Model	Predictor Response:	Estimate (95%CI)		
		Curvilinear	Straight line	Average path
Acrosome	Intercept (Independent)	0.03 (-0.15 - 0.22)	0.01 (-0.20 - 0.22)	0.01 (-0.18 - 0.19)
	Count of tracked sperm	0.70 (0.54 - 0.85)	0.60 (0.43 - 0.77)	0.74 (0.59 - 0.89)
	Morph (Satellite)	-0.09 (-0.43 - 0.25)	0.15 (-0.23 - 0.52)	0.10 (-0.24 - 0.41)
	Morph (Faeder)	-0.10 (-0.66 - 0.45)	-0.62 (-1.24 - -0.04)	-0.38 (-0.92 - 0.18)
Nucleus	Acrosome	-0.06 (-0.22 - 0.09)	0.01 (-0.16 - 0.18)	0.04 (-0.12 - 0.18)
	Intercept (Independent)	0.01 (-0.18 - 0.18)	0.01 (-0.20 - 0.22)	-0.01 (-0.19 - 0.17)
	Count of tracked sperm	0.71 (0.56 - 0.85)	0.60 (0.42 - 0.76)	0.73 (0.59 - 0.87)
	Morph (Satellite)	0.01 (-0.31 - 0.35)	0.17 (-0.21 - 0.57)	0.16 (-0.16 - 0.48)
	Morph (Faeder)	-0.10 (-0.61 - 0.42)	-0.63 (-1.23 - -0.03)	-0.37 (-0.87 - 0.12)
Midpiece	Nucleus	0.21 (0.07 - 0.36)	0.05 (-0.13 - 0.21)	0.15 (0.01 - 0.29)
	Intercept (Independent)	0.03 (-0.16 - 0.21)	0.02 (-0.20 - 0.23)	0.00 (-0.19 - 0.18)
	Count of tracked sperm	0.71 (0.56 - 0.87)	0.60 (0.42 - 0.77)	0.74 (0.58 - 0.89)
	Morph (Satellite)	-0.07 (-0.43 - 0.29)	0.15 (-0.23 - 0.54)	0.11 (-0.24 - 0.44)
	Morph (Faeder)	-0.08 (-0.64 - 0.47)	-0.62 (-1.25 - 0.03)	-0.33 (-0.89 - 0.23)
Tail	Midpiece	-0.02 (-0.18 - 0.15)	0.00 (-0.18 - 0.18)	-0.03 (-0.20 - 0.13)
	Intercept (Independent)	0.04 (-0.15 - 0.22)	0.03 (-0.17 - 0.24)	0.02 (-0.16 - 0.20)
	Count of tracked sperm	0.71 (0.56 - 0.86)	0.60 (0.43 - 0.76)	0.73 (0.58 - 0.87)
	Morph (Satellite)	-0.10 (-0.44 - 0.25)	0.11 (-0.29 - 0.50)	0.05 (-0.29 - 0.41)
	Morph (Faeder)	-0.10 (-0.65 - 0.45)	-0.64 (-1.25 - -0.04)	-0.38 (-0.91 - 0.12)
Total	Tail	0.01 (-0.15 - 0.16)	0.07 (-0.10 - 0.25)	0.07 (-0.08 - 0.22)
	Intercept (Independent)	0.04 (-0.14 - 0.23)	0.03 (-0.18 - 0.24)	0.02 (-0.16 - 0.21)
	Count of tracked sperm	0.70 (0.55 - 0.85)	0.59 (0.42 - 0.76)	0.72 (0.57 - 0.86)
	Morph (Satellite)	-0.12 (-0.46 - 0.23)	0.11 (-0.27 - 0.51)	0.05 (-0.29 - 0.39)
	Morph (Faeder)	-0.13 (-0.67 - 0.42)	-0.67 (-1.28 - -0.07)	-0.42 (-0.93 - 0.11)
Head	Total	0.07 (-0.08 - 0.22)	0.08 (-0.10 - 0.25)	0.10 (-0.05 - 0.25)
	Intercept (Independent)	0.01 (-0.17 - 0.19)	0.01 (-0.20 - 0.23)	-0.01 (-0.19 - 0.16)
	Count of tracked sperm	0.72 (0.57 - 0.87)	0.60 (0.43 - 0.76)	0.74 (0.59 - 0.88)
	Morph (Satellite)	0.00 (-0.34 - 0.35)	0.17 (-0.24 - 0.56)	0.16 (-0.16 - 0.50)
	Morph (Faeder)	-0.11 (-0.64 - 0.41)	-0.63 (-1.23 - -0.01)	-0.38 (-0.90 - 0.14)
Flagellum	Head	0.18 (0.04 - 0.33)	0.05 (-0.13 - 0.22)	0.15 (0.01 - 0.30)
	Intercept (Independent)	0.03 (-0.17 - 0.23)	0.03 (-0.18 - 0.24)	0.02 (-0.16 - 0.20)
	Count of tracked sperm	0.71 (0.56 - 0.86)	0.59 (0.42 - 0.76)	0.72 (0.58 - 0.87)
	Morph (Satellite)	-0.08 (-0.44 - 0.27)	0.10 (-0.31 - 0.50)	0.06 (-0.28 - 0.41)
	Morph (Faeder)	-0.10 (-0.68 - 0.44)	-0.66 (-1.27 - -0.05)	-0.39 (-0.91 - 0.13)
Midpiece (relative)	Flagellum	0.00 (-0.16 - 0.16)	0.07 (-0.12 - 0.24)	0.05 (-0.11 - 0.20)
	Intercept (Independent)	0.02 (-0.16 - 0.21)	0.01 (-0.20 - 0.21)	0.00 (-0.19 - 0.18)
	Count of tracked sperm	0.72 (0.57 - 0.87)	0.61 (0.44 - 0.78)	0.75 (0.61 - 0.90)
	Morph (Satellite)	-0.07 (-0.41 - 0.27)	0.16 (-0.22 - 0.54)	0.11 (-0.20 - 0.43)
	Morph (Faeder)	-0.04 (-0.60 - 0.51)	-0.58 (-1.20 - 0.05)	-0.27 (-0.81 - 0.25)
Flagellum (relative)	Midpiece (relative)	-0.06 (-0.22 - 0.10)	-0.05 (-0.23 - 0.12)	-0.10 (-0.25 - 0.05)
	Intercept (Independent)	-0.01 (-0.20 - 0.18)	0.01 (-0.20 - 0.22)	-0.02 (-0.20 - 0.17)
	Count of tracked sperm	0.73 (0.58 - 0.87)	0.59 (0.43 - 0.76)	0.74 (0.60 - 0.89)
	Morph (Satellite)	0.04 (-0.31 - 0.39)	0.15 (-0.26 - 0.56)	0.16 (-0.19 - 0.51)
	Morph (Faeder)	-0.05 (-0.59 - 0.49)	-0.63 (-1.25 - -0.01)	-0.34 (-0.85 - 0.17)
	Flagellum (relative)	-0.16 (-0.32 - 0.00)	0.00 (-0.18 - 0.18)	-0.10 (-0.25 - 0.06)

The posterior estimates (medians) of the effect sizes with the 95% credible intervals (CI) from a posterior distribution of 5,000 simulated values generated from linear models by the 'sim' function from the 'arm' R-package (Gelman & Su 2021) as presented in main text Fig. 3. Separate models were fitted for each of the three velocity-measures and for each morphological trait (in bold) while controlling for number of tracked sperm (ln-transformed) and morph. Velocity, count of tracked sperm and morphological traits were scaled (mean centered and divided by standard deviation). N = 92 males.

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