SOCIAL EVOLUTION

Genomic signatures of evolutionary transitions from solitary to group living

Karen M. Kapheim,^{1,2,3*} † Hailin Pan,^{4*} Cai Li,^{4,5} Steven L. Salzberg,^{6,7} Daniela Puiu,⁷ Tanja Magoc,⁷ Hugh M. Robertson,^{1,2} Matthew E. Hudson,^{1,8} Aarti Venkat,^{1,8,9} Brielle J. Fischman,^{1,10,11} Alvaro Hernandez,¹² Mark Yandell,^{13,14} Daniel Ence,¹³ Carson Holt,^{13,14} George D. Yocum,¹⁵ William P. Kemp,¹⁵ Jordi Bosch,¹⁶ Robert M. Waterhouse,^{17,18,19,20} Evgeny M. Zdobnov,^{17,18} Eckart Stolle,^{21,22} F. Bernhard Kraus,^{21,23} Sophie Helbing,²¹ Robin F. A. Moritz,^{21,24} Karl M. Glastad,²⁵ Brendan G. Hunt,²⁶ Michael A. D. Goodisman,²⁵ Frank Hauser,²⁷ Cornelis J. P. Grimmelikhuijzen,²⁷ Daniel Guariz Pinheiro,^{28,29} Francis Morais Franco Nunes,³⁰ Michelle Prioli Miranda Soares,²⁸ Francis Morais Franco Nunes,⁵⁰ Michelle Prion Miranda Soares,⁵⁰ Érica Donato Tanaka,³¹ Zilá Luz Paulino Simões,²⁸ Klaus Hartfelder,³² Jay D. Evans,³³ Seth M. Barribeau,³⁴ Reed M. Johnson,³⁵ Jonathan H. Massey,^{2,36} Bruce R. Southey,³⁷ Martin Hasselmann,³⁸ Daniel Hamacher,³⁸ Matthias Biewer,³⁸ Clement F. Kent,^{39,40} Amro Zayed,³⁹ Charles Blatti III,^{1,41} Saurabh Sinha,^{1,41} J. Spencer Johnston,⁴² Shawn J. Hanrahan,⁴² Sarah D. Kocher,⁴³ Jun Wang,^{4,44,45,46,47}† Gene E. Robinson,^{1,48}† Guojie Zhang^{4,49}†

The evolution of eusociality is one of the major transitions in evolution, but the underlying genomic changes are unknown. We compared the genomes of 10 bee species that vary in social complexity, representing multiple independent transitions in social evolution, and report three major findings. First, many important genes show evidence of neutral evolution as a consequence of relaxed selection with increasing social complexity. Second, there is no single road map to eusociality; independent evolutionary transitions in sociality have independent genetic underpinnings. Third, though clearly independent in detail, these transitions do have similar general features, including an increase in constrained protein evolution accompanied by increases in the potential for gene regulation and decreases in diversity and abundance of transposable elements. Eusociality may arise through different mechanisms each time, but would likely always involve an increase in the complexity of gene networks.

he evolution of eusociality involves changes

in the unit of natural selection, from the

individual to a group (1). Bees evolved

eusociality multiple times and are ex-

tremely socially diverse (2) (Fig. 1), but all

pollinate angiosperms, including many crops

essential to the human diet (3). Simple euso-

ciality may be facultative or obligate, and both

forms are characterized by small colonies with

a reproductive queen and one or more workers

that, due to social and nutritional cues, forego

reproduction to cooperatively care for their

siblings (2). Further evolutionary elaborations

have led to complex eusociality, "superorgan-

isms" with colonies of several thousand individ-

uals, sophisticated modes of communication,

and morphological specializations for division

Theory predicts that the evolution of simple

eusociality involves increased regulatory flex-

ibility of ancestral gene networks to create

specialized reproductive and nonreproductive

individuals, and the evolution of complex eu-

sociality requires genetic novelty to coordinate

emergent properties of group dynamics (5). To

test these predictions, we analyzed five de novo

and five publicly available draft genome sequences of 10 bee species from three families, representing two independent origins of eusociality in Apidae and Halictidae and two independent elaborations of simple to complex eusociality in two apid tribes [Apini (honeybees) and Meliponini (stingless bees); Fig. 1]. The draft genomes were of comparable, high quality (sup-We found that the transition from solitary to

group life is associated with an increased capacity for gene regulation. We scanned the promoter regions of 5865 single-copy orthologs among the 10 species to calculate a motif score [representing the number and binding strength of experimentally characterized transcription factor binding sites (TFBSs)] for 188 Drosophila melanogaster TFs (6) with at least one ortholog in each of the 10 bees, and correlated motif score with social complexity, using phylogenetically independent contrasts (7). Of 2101 significantly correlated motif-gene pairs, 89% were positive and 11% negative, showing that TFs tend to have increased capacity to regulate genes in eusocial species of bees, relative to solitary species (Fig. 2A, supplementary materials).

plementary materials).

Further evidence for increased capacity for gene regulation throughout social evolution is a positive ranked correlation between social complexity and the number of genes predicted to be methylated (7) (Spearman's rho = 0.76, P = 0.01; phylogenetically corrected Spearman's rho = 0.64, P = 0.06; Fig. 2B; bioinformatics predictions validated with bisulfite sequencing data for three invertebrate species; supplementary materials). DNA methylation affects gene expression in a variety of ways (8). Thus, this result suggests an expansion in regulatory capacity with increasingly sophisticated sociality.

The potential for increased regulatory capacity was further revealed at the protein-coding level. Increased social complexity also is associated with rapid evolution of genes involved in coordinating gene regulation. A Bayesian phylogenetic covariance analysis (9) of 5865 single-copy orthologs identified 162 genes with accelerated evolution in species with increased social complexity (7) (additional data table S3). These rapidly evolving genes were significantly enriched (P < 0.05) for Gene Ontology (GO) terms related to regulation of transcription, RNA splicing, ribosomal structure, and regulation of translation (supplementary text and tables S11 and S12). Similar results have been reported for bee and ant species (10-13); our findings reveal the underlying causes. Approximately two-thirds of these genes are under stronger directional selection in species with increasingly complex eusociality, but we also detected nonadaptive evolution. One-third of the rapidly evolving genes are under relaxed purifying selection in species with complex eusociality, possibly due to reduced effective population sizes (14).

We also found an additional 109 genes, significantly enriched (P < 0.05) for functions related to protein transport and neurogenesis, which evolve slower with increased social complexity (supplementary text, table S13, and additional data table S3). This includes orthologs of derailed 2 and *frizzled*, which function as Wnt signaling receptors in Drosophila synaptogenesis (15), and rigor mortis, a nuclear receptor involved in hormone signaling (16). A similar pattern of reduced evolutionary rate has been described for genes expressed in human and honey bee brains, potentially due to increasing pleiotropic constraint in complex gene networks (17, 18). Constrained protein evolution of neural and endocrine-related genes seems at odds with the evolution of complexity, but this constraint appears to be compensated for, or perhaps driven by, increased capacity for gene regulation.

We next investigated whether these molecular evolution patterns involve similar sets of genes and cis-regulatory elements among the early (facultative and obligate simple eusociality) and advanced (complex eusociality) stages of independent social transitions. We identified lineage-specific differences in coding sequences and promoter regions of 1526 "social genes" for which evolutionary rate (dN/dS) is faster or slower with increased social complexity in two independent origins and two independent elaborations of eusociality (7)

of labor (4).

(Fig. 1). Among these lineage-specific social genes, we found common patterns of cis-regulatory evolution: gains of TFBSs in the promoters of genes that evolve slower with increasing social complexity (Fig. 2C and supplementary text). This suggests that a shared feature of both independent origins and elaborations of eusociality is increasingly constrained protein evolution with increasing potential for novel gene expression patterns. The TFs responsible for this pattern were different for each social transition, even though our analysis was limited to highly conserved TFs (Table 1). Several function in neurogenesis or neural plasticity, or are prominent regulators of endocrine-mediated brain gene expression in honeybees (19, 20).

We found further lineage-specific differences among the rapidly evolving "social genes" themselves. Genes undergoing accelerated evolution at the origins of eusociality were significantly enriched for GO terms related to signal transduction in both Apidae and Halictidae, but they shared only six genes (6 out of 354 and 167 genes, respectively; hypergeometric test, P = 0.82; Fig. 2D and additional data tables S5 and S6). Rapid evolution of signal transduction pathways may be a necessary step in all origins of eusociality to mediate intracellular responses to novel social and environmental stimuli (10), but selection appears to have targeted different parts of these pathways in each independent transition. Castespecific expression and other analyses of these genes are needed to determine their function in eusociality.

Genes showing signatures of rapid evolution with the elaborations of complex eusociality were also highly disparate between honeybees and stingless bees, with only 43 shared genes and no shared enriched GO terms (43 out of 625 and 512 genes, respectively; hypergeometric test, P =0.70; Fig. 2D and additional data tables S5 and S6). In addition, only 2 out of 5865 single-copy orthologs showed a signature of convergent evolution by fitting a dendrogram based on social complexity significantly better than the accepted molecular phylogeny (7) (supplementary text and fig. S21). Similarly, families of major royal jelly protein genes, sex-determining genes, odorant receptors, and genes involved in lipid metabolism expanded in some, but not all, lineages of complex eusocial bees (7) (Table 2 and supplementary text). These results suggest that gene family expansion is associated with complex eusociality as predicted (*5*), but involves different genes in each case. Despite striking convergence of social traits among the superorganisms (*4*), the final stages of transformation to this level of biological organization do not necessarily involve common molecular pathways.



Fig. 1. Phylogeny and divergence times (28) of bees selected for genome analysis. We analyzed two independent origins of simple eusociality from a solitary ancestor, one each in Apidae (white circle 1) and Halictidae (white circle 2), and two independent elaborations of complex eusociality in honeybees (gray circle 1) and stingless bees (gray circle 2). Most bees mate once, but honeybees mate with multiple males. All bees eat pollen and nectar from flowering plants. Species names are colored according to degree of social complexity: blue: ancestrally solitary; green: facultative simple eusociality; orange: obligate simple eusociality; red: obligate complex eusociality. The social biology of *E. mexicana* is unknown, but is representative of the facultative simple eusocial life history (*29*). Numbers in each box are approximate colony size on a log scale. MRCA, most recent common ancestor; mya, millions of years ago.

Downloaded from http://science.sciencemag.org/ on September 21, 2019

1Carl R. Woese Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ²Department of Entomology, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ³Department of Biology, Utah State University, Logan, UT 84322, USA. ⁴China National GeneBank, BGI-Shenzhen, Shenzhen, 518083, China. ⁵Centre for GeoGenetics, Natural History Museum of Denmark, University of Copenhagen, Copenhagen, 1350, Denmark. ⁶Departments of Biomedical Engineering, Computer Science, and Biostatistics, Johns Hopkins University, Baltimore, MD 21218, USA. ⁷Center for Computational Biology, McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine, Baltimore, MD 21205, USA. ⁸Department of Crop Sciences, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ⁹Department of Human Genetics, University of Chicago, Chicago, IL 60637, USA. ¹⁰Program in Ecology and Evolutionary Biology, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ¹¹Department of Biology, Hobart and William Smith Colleges, Geneva, NY 14456, USA. ¹²Roy J. Carver Biotechnology Center, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ¹³Department of Human Genetics, Eccles Institute of Human Genetics, University of Utah, Salt Lake City, UT 84112, USA. ¹⁴USTAR Center for Genetic Discovery, University of Utah, Salt Lake City, UT 84112, USA. ¹⁵U.S. Department of Agriculture-Agricultural Research Service (USDA-ARS) Red River Valley Agricultural Research Center, Biosciences Research Laboratory, Fargo, ND 58102, USA. ¹⁶Center for Ecological Research and Forestry Applications (CREAF), Universitat Autonoma de Barcelona, 08193 Bellaterra, Spain. ¹⁹Department of Genetic Medicine and Development, University of Geneva Medical School, 1211 Geneva, Switzerland. ¹⁸Swiss Institute of Bioinformatics, 1211 Geneva, Switzerland. ¹⁹Computer Science and Artificial Intelligence Laboratory, Massachusetts Institute of Technology (MIT), Cambridge, MA 02139, USA. ²⁰The Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA. ²¹Institute of Biology, Department Zoology, Martin-Luther-University Halle-Wittenberg, Hoher Weg 4, D-06099 Halle (Saale), Germany. ²²Queen Mary University of London, School of Biological and Chemical Sciences Organismal Biology Research Group, London E1 4NS, UK. ²³Department of Laboratory Medicine, University Hospital Halle, Ernst Grube Strasse 40, D-06120 Halle (Saale), Germany.²⁴German Centre for Integrative Biodiversity Research (iDiv), Halle-Jena-Leipzig, 04103 Leipzig, Germany. ²⁵School of Biology, Georgia Institute of Technology, Atlanta, GA 30332, USA. ²⁶Department of Entomology, University of Georgia, Griffin, GA 30223, USA. ²⁷Center for Functional and Comparative Insect Genomics, Department of Biology, University of Copenhagen, Copenhagen, Denmark. 28 Departamento de Biologia, Faculdade de Filosofia, Ciências e Letras de Ribeirão Preto, Universidade de São Paulo, 14040-901 Ribeirão Preto, SP, Brazil.²⁹Departamento de Tecnologia, Faculdade de Ciências Agrárias e Veterinárias, Universidade Estadual Paulista (UNESP), 14884-900 Jaboticabal, SP, Brazil. 30 Departamento de Genética e Evolução, Centro de Ciências Biológicas e da Saúde, Universidade Federal de São Carlos, 13565-905 São Carlos, SP, Brazil. ³¹ Departamento de Genética, Faculdade de Medicina de Ribeirão Preto, Universidade de São Paulo, 14049-900 Ribeirão Preto, SP, Brazil.³²Departamento de Biología Celular e Molecular e Bioagentes Patogênicos, Faculdade de Medicina de Ribeirão Preto, Universidade de São Paulo, 14049-900 Ribeirão Preto, SP, Brazil.³³USDA-ARS Bee Research Lab, Beltsville, MD 20705 USA.³⁴Department of Biology, East Carolina University, Greenville, NC 27858, USA. 35Department of Entomology, Ohio Agricultural Research and Development Center, Ohio State University, Wooster, OH 44691, USA. 36Department of Ecology and Evolutionary Biology, University of Michigan, Ann Arbor, MI 48109, USA. ³⁷Department of Animal Sciences, University of Illinois, Urbana, IL 61801, USA. ³⁸Department of Population Genomics, Institute of Animal Husbandry and Animal Breeding, University of Hohenheim, Germany. ³⁹Department of Biology, York University, Toronto, ON M3J 1P3, Canada. ⁴⁰Janelia Farm Research Campus, Howard Hughes Medical Institue, Ashburn, VA ⁴³Department of Organismic and Evolutionary Biology, Museum of Comparative Zoology, Harvard University, Cambridge, MA 02138, USA. ⁴⁴Department of Biology, University of Copenhagen, 2200 Denmark. ⁴⁵Princess AI Jawhara Center of Excellence in the Research of Hereditary Disorders, King Abdulaziz University, Jeddah 21589, Saudi Arabia. ⁴⁶Macau University of Science and Technology, Avenida Wai long, Taipa, Macau 999078, China. ⁴⁷Department of Medicine, University of Hong Kong, Hong Kong, Hong Kong, de Center for Advanced Study Professor in Entomology and Neuroscience, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA. ⁴⁹Centre for Social Evolution, Department of Biology, Universitetsparken 15, University of Copenhagen, DK-2100 Copenhagen, Denmark. *These authors contributed equally to this work. +Corresponding author. E-mail: karen.kapheim@usu.edu (K.M.K.); wangj@genomics.org.cn (J.W.); generobi@illinois.edu (G.E.R.); zhanggj@genomics.org.cn (G.Z.)





levels of social complexity [from (D)]. Background shading follows circle shading in Fig. 1. (**D**) Number of genes for which evolutionary rate is faster or slower in lineages with higher compared to lower social complexity. Pie charts represent the proportion of genes evolving slower (light green) or faster (dark orange) with increased social complexity. Venn diagram shading follows circle shading in Fig. 1. (**E**) Complex eusocial species have a reduced proportion of repetitive DNA compared to other bees (see text for statistics). LTR, long terminal repeat; LINE, long interspersed element; SINE, short interspersed element; DNA, DNA transposon; LARD, large retrotransposon derivative; TRIM, terminal repeat retrotransposon in miniature; MITE, miniature inverted-repeat transposable element; TES, transposable elements.

The major transitions in evolution involve a reduction in conflict as the level of natural selection rises from the individual to the group (*I*). Extending this to intragenomic conflict may explain our finding of decreased diversity and abundance of transposable elements (TEs) with increasing social complexity (7) (regression after phylogenetic correction, F = 8.99, adjusted $R^2 = 0.47$, P = 0.017; Fig. 2E, figs. S42 to S44, and supplementary text). This may be a consequence of increased recombination rates among highly eusocial insects (21, 22) or because key features of

complex eusociality lead to decreased exposure to parasites and pathogens that horizontally transmit TEs (4, 23). Eusociality in bees may thus provide natural immunity against certain types of intragenomic conflict.

Our results and those in (10–13) support the prediction that changes in gene regulation are key features of evolutionary transitions in biological organization (5). Our results further reveal the convergent adaptive and nonadaptive evolutionary processes common to both the early and advanced stages of multiple inde-

Table 1. Transcription factors (TFs) and corresponding motifs associated with origins and elaborations of eusociality in bees. [Motif names: Fly Factor Survey (6); supplementary text.]

Motif	D. melanogaster TFs	Hypergeometric test <i>P</i> -value	
	Solitary to simple eusociality–Apidae		
lola_PQ_SOLEXA	Lola	0.0047	
	Solitary to simple eusociality–Halictidae		
br_PL_SOLEXA_5	Br	0.0016	
S	<i>Simple eusociality to complex eusociality–honeybees</i>		
h_SOLEXA_5	dpn,h	0.0027	
Simple eusociality to complex eusociality–stingless bees			
Side_SOLEXA_5	E_spl, HLHm3, HLHm5,		
	HLHm7, HLHmbeta,		
	HLHmdelta, HLHmgamma, Side	0.0008	
usp_SOLEXA	EcR,svp,usp	0.0013	
CrebA_SOLEXA	CrebA	0.0040	
CG5180_SOLEXA	CG5180	0.0044	
tai_Met_SOLEXA_5	5 Mio_bigmax,tai_Met	0.0045	
ttk_PA_SOLEXA_5	Ttk	0.0078	
gsb_SOLEXA	gsb,Poxn,prd	0.0083	
tai_SOLEXA_5	Tai	0.0100	

Table 2. Relative size of select gene families as related to social complexity in bees.

Family	Function	Eusocial bees compared to solitary bees
	Differences among bees	
Major royal jelly	Brood feeding	Expanded only in Apis
Sex determination pathway genes	Sex-specific development	Expanded in some eusocial lineages
Odorant receptors	Olfaction	Expanded in complex eusocial lineages
	Metabolic processing of	Expanded in complex
Lipid metabolism genes	lipids	eusocial lineages
	Similarities across bees	
Biogenic amines receptors, neuropeptides, GPCRs*	Neural plasticity	Similar
Insulin-signaling and	Insect development, caste	Similar
ecdysone pathway genes	determination in honeybees, behavioral plasticity as adults	
Immunity	Infectious disease protection	Similar
Cytochrome P450	Detoxification	Similar
monooxygenase genes		

*GPCRs, G protein-coupled receptors.

It is now clear that there are lineage-specific genetic changes associated with independent origins of eusociality in bees, and independent elaborations of eusociality in both bees and ants. This includes different sets of genes showing caste-biased expression across species (24–26) and, as we have shown, evolutionary modifications of TEs, gene methylation, and cis-regulatory patterns associated with the suite of life-history traits that define eusociality. This suggests that if it were possible to "replay life's tape" (27), eusociality may arise through different mechanisms each time, but would likely always involve an increase in the complexity of gene networks.

pendent transitions from solitary to group living.

REFERENCES AND NOTES

- J. Maynard Smith, E. Szathmáry, *The Major Transitions in Evolution* (Oxford Univ. Press, Oxford, UK, 1995).
- C. D. Michener, *The Social Behavior of the Bees* (Harvard Univ. Press, Cambridge, MA, 1974).
- A.-M. Klein et al., Proc. Biol. Sci. B 274, 303–313 (2007).
 H. Hölldobler, E. O. Wilson, The Superorganism: The Beauty, Elegance and Strangeness of Insect Societies (Norton,
- New York, 2009). 5. B. R. Johnson, T. A. Linksvayer, *Q. Rev. Biol.* **85**, 57–79
- (2010).L. J. Zhu et al., Nucleic Acids Res. 39, D111–D117 (2011).Materials and methods are available as supplementary
- materials on Science Online. 8. H. Yan et al., Annu. Rev. Entomol. **60**, 435–452 (2015).
- 9. N. Lartillot, R. Poujol, *Mol. Biol. Evol.* **28**, 729–744 (2011).
- 10. S. H. Woodard et al., Proc. Natl. Acad. Sci. U.S.A. 108, 7472–7477 (2011).
- 11. B. A. Harpur et al., Proc. Natl. Acad. Sci. U.S.A. 111, 2614–2619 (2014).
- 12. J. Roux et al., Mol. Biol. Evol. 31, 1661-1685 (2014).
- 13. D. F. Simola et al., Genome Res. 23, 1235-1247 (2013)
- 14. J. Romiguier et al., J. Evol. Biol. 27, 593-603 (2014).
- 15. M. Park, K. Shen, EMBO J. 31, 2697-2704 (2012).
- J. Gates, G. Lam, J. A. Ortiz, R. Losson, C. S. Thummel, Development 131, 25–36 (2004).
- D. Brawand *et al.*, *Nature* **478**, 343–348 (2011).
 D. Molodtsova, B. A. Harpur, C. F. Kent, K. Seevananthan,
- A. Zayed, Front. Genet. 5, 431 (2014).
- D. W. Pfaff, A. P. Arnold, A. M. Etgen, R. T. Rubin, S. E. Fahrbach, Eds., *Hormones, Brain and Behavior* (Elsevier, New York, 2009).
- S. Chandrasekaran *et al.*, *Proc. Natl. Acad. Sci. U.S.A.* 108, 18020–18025 (2011).
- 21. L. Wilfert, J. Gadau, P. Schmid-Hempel, *Heredity* **98**, 189–197 (2007).
- E. S. Dolgin, B. Charlesworth, *Genetics* **178**, 2169–2177 (2008).
- S. Schaack, C. Gilbert, C. Feschotte, *Trends Ecol. Evol.* 25, 537–546 (2010).
- 24. B. Feldmeyer, D. Elsner, S. Foitzik, *Mol. Ecol.* 23, 151–161 (2014).
- P. G. Ferreira et al., Genome Biol. 14, R20 (2013).
 B. G. Hunt et al., Proc. Natl. Acad. Sci. U.S.A. 108,
- 15936–15941 (2011). 27. S. J. Gould, Wonderful Life: The Burgess Shale and the Nature
- of History (Norton, New York, 1989).
 28. S. Cardinal, B. N. Danforth, Proc. Biol. Sci. 280, 20122686 (2013).
- (2013).
 S. Cardinal, B. N. Danforth, *PLOS ONE* 6, e21086 (2011).

ACKNOWLEDGMENTS

Data deposition at National Center for Biotechnology Information: *H. laboriosa, D. novaeangliae, E. mexicana, M. quadrifasciata,* and *M. rotundata* genome assemblies accession numbers PRJNA279436, PRJNA279825, PRJNA279814, PRJNA279820, and PRJNA66515. Funding for genome sequencing and analysis: BGI, a U.S. National Institutes of Health Pioneer Award (DP1 OD006416) to G.E.R., and European Union Marie Curie International Incoming Fellowship (300837) to G.Z. Additional

funding: U.S. National Science Foundation grant DEB-0640690 (M.A.D.G.) and DEB-0743154 (G.E.R. and M.E.H); Danish Council for Independent Research grants 10-081390 (C.L.) and 0602-01170B (C.J.P.G.); Lundbeck Foundation (C.J.P.G); Georgia Tech-Elizabeth Smithgall Watts endowment (M.A.D.G.); Marie Currie International Outgoing Fellowship PIOF-GA-2011-303312 (R.M.W.); Swiss National Science Foundation award 31003A-125350, Commission Informatique of the University of Geneva, Schmidheiny Foundation, and Swiss Institute of Bioinformatics (E.M.Z.); and Natural Sciences and Engineering Research Council of Canada Discovery Grant and Early Research Award (A.Z.). This project was conducted under the auspices of the i5K Initiative. We thank the Roy J. Carver Biotechnology Center (sequencing services); N. Lartillot (advice on Coevol); T. Newman (assistance with DNA extractions); and E. Hadley (assistance with figures). Computational support: D. Davidson, N. Band, D. Slater (University of Illinois), J. H. Kidner and H. Scharpenberg (Martin-Luther-University Halle-Wittenberg); Compute Canada; and Center for High Performance Computing at the University of Utah. J. Himes created the illustrations in Fig. 1. We are grateful to T. Pitts-Singer, H. G. Hall, B. N. Danforth, J. Gibbs, and S. Cardinal for providing bee specimens; R. Ayala for identification of *E. mexicana*; J. Vega and the staff at Estación de Biología Chamela, Universidad Nacional Autónorma de México (UNAM), for support during collecting trips; and S. A. Cameron for insightful discussion.

HUMAN OOCYTES

Error-prone chromosome-mediated spindle assembly favors chromosome segregation defects in human oocytes

Zuzana Holubcová,¹ Martyn Blayney,² Kay Elder,² Melina Schuh¹*

Aneuploidy in human eggs is the leading cause of pregnancy loss and several genetic disorders such as Down syndrome. Most aneuploidy results from chromosome segregation errors during the meiotic divisions of an oocyte, the egg's progenitor cell. The basis for particularly error-prone chromosome segregation in human oocytes is not known. We analyzed meiosis in more than 100 live human oocytes and identified an error-prone chromosome segregation defects. Human oocytes assembled a meiotic spindle independently of either centrosomes or other microtubule organizing centers. Instead, spindle assembly was mediated by chromosomes and the small guanosine triphosphatase Ran in a process requiring ~16 hours. This unusually long spindle assembly period was marked by intrinsic spindle instability and abnormal kinetochore-microtubule attachments, which favor chromosome segregation errors and provide a possible explanation for high rates of aneuploidy in human eggs.

eiosis in human oocytes is more prone to chromosome segregation errors than mitosis (1, 2), meiosis during spermatogenesis (3, 4), and female meiosis in other organisms (3, 5). Despite its importance for fertility and human development, meiosis in human eggs has hardly been studied. Human oocytes are only available in small numbers, warranting single-cell assays capable of extracting maximal information. Although highresolution live-cell microscopy is an ideal method, oocyte development in the ovary poses challenges to direct imaging. We therefore established an experimental system (6) for ex vivo high-resolution fluorescence microscopy of human oocytes freshly harvested from women undergoing gonadotropin-stimulated in vitro fertilization cycles. To establish the major stages of meiosis in this system, we simultaneously monitored microtubules and chromosomes for ~24 to 48 hours (Fig. 1 and movie S1). Similar to the situation in situ (7), human oocytes matured into fertilizable eggs over this time course, as judged

by the formation of a polar body. The morphologically identifiable stages (Fig. 1A) at characteristic times after nuclear envelope breakdown [(NEBD), set to 0 hours] provided a time-resolved framework for human oocyte meiosis (Fig. 1B). This reference timeline post-NEBD is used throughout this paper.

Before NEBD, chromosomes were highly condensed and clustered around the nucleolus. Instead of rapidly nucleating microtubules upon NEBD, human oocytes first formed a chromosome aggregate that was largely devoid of microtubules (Fig. 1A; movie S1; and fig. S1, A and B). Microtubules were first observed at ~5 hours, when they started to form a small aster within the chromosome aggregate. As the microtubule aster grew, the chromosomes became individualized and oriented on the surface of the aster with their kinetochores facing inwards. The microtubule aster then extended into an early bipolar spindle that carried the chromosomes on its surface (Fig. 1A; movie S1; and fig. S1, C to E). The chromosomes then entered the spindle but remained distributed throughout the entire spindle volume. Chromosomes first congressed in the spindle center at ~13 hours but continued to oscillate around the spindle equator. Stable chromosome alignment was typically only achieved

SUPPLEMENTARY MATERIALS

www.sciencemag.org/content/348/6239/1139/suppl/DC1 Materials and Methods Supplementary Text Figs. S1 to S44 Tables S1 to S32 Additional Data Tables S1 to S12 References (*30–151*) Author Contributions

19 December 2014; accepted 6 May 2015 Published online 14 May 2015; 10.1126/science.aaa4788

close to anaphase onset (Fig. 1, A and B, and movie S1). Unexpectedly, the spindle volume increased over the entire course of meiosis, up until anaphase onset (Fig. 1, C and D). The barrelshaped spindle formed in this process consisted of loosely clustered bundles of microtubules and lacked astral microtubules (movie S2 and fig. S2). At ~17 hours, the oocytes progressed into anaphase and eliminated half of the homologous chromosomes in a polar body. Nearly a day after NEBD, the oocytes had formed a bipolar metaphase II spindle and matured into a fertilizable egg. The stages and timing of meiosis were highly reproducible among oocytes (Fig. 1, A and B) and could also be observed in fixed oocytes (fig. S1, A to I). Importantly, 79.0% of imaged human oocytes extruded a polar body. This indicates that the imaging assays, as well as the methods by which the oocytes were obtained and processed, did not have a prominent effect on meiotic progression.

The surprisingly slow and gradual build-up of the spindle over 16 hours (Fig. 1, C and D) is in stark contrast to mitosis, where spindle assembly takes only ~30 min (8), or meiosis in mouse oocytes, where it takes 3 to 5 hours (9-11). During mitosis, two centrosomes ensure the rapid assembly of a spindle. In oocytes of many species, centrosomes are absent but functionally replaced by microtubule organizing centers (MTOCs) that lack centrioles (9, 12). Human oocvtes also lack centrosomes (13-15), but whether acentriolar MTOCs participate in spindle assembly is unclear (16-19). We consistently detected pericentrinand γ -tubulin-positive MTOCs at the spindle poles of mitotic cells and metaphase I and II (MI and MII) mouse oocytes, but never at MI or MII spindles in human oocytes (Fig. 2, A and B, and fig. S3). Thus, our data suggest that meiotic spindles in human oocytes lack detectable MTOCs.

In *Xenopus* egg extracts, chromosomes can serve as sites of microtubule nucleation if centrosomes are absent (20). The human oocytes we imaged also initiated microtubule nucleation in the region of the chromosome aggregate (78 of 78 live human oocytes). High-resolution imaging of fixed human oocytes confirmed that microtubules were first nucleated on chromosomes, emanating primarily from kinetochores (Fig. 2C, movie S3, and fig. S4). MTOC-nucleated cytoplasmic asters, such as those seen in chromosomal proximity upon NEBD in mouse oocytes (9), could not be detected. Thus, chromosomes, not MTOCs, serve as major sites of microtubule nucleation in human oocytes.

¹Medical Research Council, Laboratory of Molecular Biology, Francis Crick Avenue, Cambridge Biomedical Campus, Cambridge CB2 0QH, UK. ²Bourn Hall Clinic, Bourn, Cambridge CB23 2TN, UK.

^{*}Corresponding author. E-mail: mschuh@mrc-lmb.cam.ac.uk

Science

Genomic signatures of evolutionary transitions from solitary to group living

Karen M. Kapheim, Hailin Pan, Cai Li, Steven L. Salzberg, Daniela Puiu, Tanja Magoc, Hugh M. Robertson, Matthew E. Hudson, Aarti Venkat, Brielle J. Fischman, Alvaro Hernandez, Mark Yandell, Daniel Ence, Carson Holt, George D. Yocum, William P. Kemp, Jordi Bosch, Robert M. Waterhouse, Evgeny M. Zdobnov, Eckart Stolle, F. Bernhard Kraus, Sophie Helbing, Robin F. A. Moritz, Karl M. Glastad, Brendan G. Hunt, Michael A. D. Goodisman, Frank Hauser, Cornelis J. P. Grimmelikhuijzen, Daniel Guariz Pinheiro, Francis Morais Franco Nunes, Michelle Prioli Miranda Soares, Érica Donato Tanaka, Zilá Luz Paulino Simões, Klaus Hartfelder, Jay D. Evans, Seth M. Barribeau, Reed M. Johnson, Jonathan H. Massey, Bruce R. Southey, Martin Hasselmann, Daniel Hamacher, Matthias Biewer, Clement F. Kent, Amro Zayed, Charles Blatti III, Saurabh Sinha, J. Spencer Johnston, Shawn J. Hanrahan, Sarah D. Kocher, Jun Wang, Gene E. Robinson and Guojie Zhang

Science **348** (6239), 1139-1143. DOI: 10.1126/science.aaa4788originally published online May 14, 2015

For bees, many roads lead to social harmony

Eusociality, where workers sacrifice their reproductive rights to support the colony, has evolved repeatedly and represents the most evolved form of social evolution in insects. Kapheim *et al.* looked across the genomes of 10 bee species with varying degrees of sociality to determine the underlying genomic contributions. No one genomic path led to eusociality, but similarities across genomes were seen in features such as increases in gene regulation and methylation. It also seems that selection pressures relaxed after the emergence of complex sociality. *Science*, this issue p. 1139

ARTICLE TOOLS	http://science.sciencemag.org/content/348/6239/1139
SUPPLEMENTARY MATERIALS	http://science.sciencemag.org/content/suppl/2015/05/13/science.aaa4788.DC1
REFERENCES	This article cites 138 articles, 27 of which you can access for free http://science.sciencemag.org/content/348/6239/1139#BIBL
PERMISSIONS	http://www.sciencemag.org/help/reprints-and-permissions

Use of this article is subject to the Terms of Service

Science (print ISSN 0036-8075; online ISSN 1095-9203) is published by the American Association for the Advancement of Science, 1200 New York Avenue NW, Washington, DC 20005. The title *Science* is a registered trademark of AAAS.

Copyright © 2015, American Association for the Advancement of Science