Two-Parameter Characterization of Chromosome-Scale Recombination Rate

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ABSTRACT:
The genome-wide recombination rate ($R_R$) of a species is often described by one parameter, the ratio between total genetic map length ($G$) and physical map length ($P$), measured in centiMorgans per Megabase (cM/Mb). The value of this parameter varies greatly between species, but the cause for these differences is not entirely clear. A constraining factor of overall $R_R$ in a species, which may cause increased $R_R$ for smaller chromosomes, is the requirement of at least one chiasma per chromosome (or chromosome-arm) per meiosis. In the present study, we quantify the relative excess of recombination events on smaller chromosomes by a linear regression model, which relates the genetic length of chromosomes to their physical length. We find for several species that the two-parameter regression, $G = G_0 + k \cdot P$ provides a better characterization of the relationship between genetic and physical map length than the one-parameter regression that runs through the origin. A non-zero intercept ($G_0$) indicates a relative excess of recombination on smaller chromosomes in a genome. Given $G_0$, the parameter $k$ predicts the increase of genetic map length over the increase of physical map length. The observed values of $G_0$ have a similar magnitude for diverse species, whereas $k$ varies by two orders of magnitude. The implications of this strategy for the genetic maps of human, mouse, rat, chicken, honeybee, worm and yeast are discussed.
Introduction

The rate of meiotic recombination rate (RR), defined as the ratio between genetic and physical map length and measured in centiMorgan per Megabase (cM/Mb), is known to vary widely between the genomes of different species. As a rule of thumb for the human genome, 1cM genetic map length equals 1Mb physical map length, see e.g. (Collins and Morton 1998; Ulgen and Li 2005). This rate is about twice as large as the genome-wide RR observed in the mouse genome (Jense-Seaman et al. 2004), but far less than the RR of 340cM/Mb that is observed in the yeast genome (Mortimer et al. 1992; Baudat and Nicolas 1997). Understanding of these differences in RR between different species is of fundamental importance for evolutionary and medical genetics (Nachman 2002). In addition to these differences between species, it was also noted that RR differs between chromosomes within a species, with smaller chromosomes showing higher RR (Nachman and Churchill 1996; Broman et al. 1998; Lander et al. 2001; Venter et al. 2001; Kong et al. 2002; Matise et al. 2007). Therefore, species differences in genome-wide RR may be best studied under a model that also considers the intragenomic differences between chromosomes.

From a population genetic perspective, the main role of recombination is the production of new combinations of alleles by shuffling of parental haplotypes, which increases the efficiency of natural selection in theoretical and empirical model systems (Maynard-Smith 1978; Barton and Charlesworth 1998; Rice 2002; Otto and Lenormand 2002). Many recent empirical studies have addressed the question at which sites in a genome recombination is most likely to occur (Petes 2001; McVean et al. 2004; Hey 2004; Myers et al. 2005; Coop 2005; Mancera et al. 2008). In this context it was also found that RR evolves extremely fast on a kb-scale (Ptak et al. 2005; Winckler et al. 2005) and that historical recombination hotspots are associated with specific gene functions in human, which was hypothesized to indicate an influence of natural selection on hotspot locations (Freudenberg et al. 2007; The International HapMap Consortium 2007). When RR is examined at a megabase scale instead of a kilobase scale, the evolution of local RR is more constrained (Myers et al. 2005), and differs much less between closely related species, such as human and chimpanzees (Winckler et al. 2005; Ptak et al. 2005). However, the mechanism behind this conservation of RR on the larger
scale is unclear. One contributing explanation could be the requirement of a minimal or fixed number of chiasmata per chromosome during meiosis to stabilize homologous chromosome pairs (Mather 1938).

The question how many chiasmata are exactly required per chromosome or per chromosome arm has not been resolved yet and might not have a generally valid answer (Lynn et al. 2004; Laurie and Hultén 1985). Nevertheless, these meiotic constraints can explain the excess of recombination on shorter chromosomes. Consistent with an influence of karyotype on overall recombination rate, a correlation was found between the number of chromosome arms in a genome and the genetic map length (De Villena and Sapienza 2001a). Altered recombination may lead to aneuploidy (Hassold and Hunt 2001; Lynn et al. 2004), which may impose strong selective constraints and explain the tight relationship between karyotype structure and recombination rate (Dumas and Britton-Davidian 2002; De Villena and Sapienza 2001b).

On the other hand, domesticated plants and animals show evidence for increased chiasma formation (Burt and Bell 1987), which suggests that there exist additional determinants of genome-wide RR than karyotype. For instance, the level of interference in chiasma formation could differ between species (Broman et al. 2002). Therefore, it would be useful to apply a formal method that separates the contribution of karyotype structure from the relationship between physical and genetic map length. This is not accomplished by the genome-wide cM/Mb ratio: although the cM/Mb ratio is a convenient single parameter measurement, it does not model the higher contribution of smaller chromosomes to the genome-wide RR of a species.

To address this problem and better understand the overall RR of a genome, we propose a novel strategy that explicitly models, if and to what extent, the overall RR in a genome is influenced by the relative excess of recombination on smaller chromosomes. This proposed two-parameter strategy takes into account that a certain minimal amount of recombination is required to maintain genome integrity during meiosis (Mather 1938) and that a genome therefore has minimal genetic map length. This idea becomes more clear, if we use a statistical regression framework to compare the proposed strategy with the one parameter strategy that is typically applied to shorter scales than the chromosome-scale. Since the one-parameter characterization of RR implies that genetic length is proportional to the physical length and
recombination events occur independently on different chromosomes, the cM/Mb ratio is the slope of the linear regression of genetic lengths of chromosomes over their sequence lengths, with the requirement that the regression line goes through the origin. In our new approach, we drop the requirement that the regression line must go through origin by using two parameters to fit the genome-wide genetic map information at the chromosomal scale.

From a biological perspective, the one-parameter model considers the length of the genetic map of a genome to be determined by the length of the underlying physical map and the species-specific $RR$. Building on this, the two-parameter model also includes a separate effect of karyotype structure that may produce a disproportional distribution of recombination events over chromosomes of different length. Under the two-parameter model, the value of the $y$-intercept quantifies the relative excess of recombination events on a hypothetical chromosome with length zero, whereas the slope of the regression measures the increase of genetic with physical map length in the same way as the one-parameter model. Our results show that in human, as well as other species, the two-parameter regression provides a much better fit for describing the genetic map length of chromosomes.

Results

A two-parameter regression model fits the genetic map length of human chromosomes better than the one-parameter model

To look for systematic differences in recombination rate between human chromosomes, we started by reproducing the Marey map (Chakravarti 1991; Rezvoy et al. 2007), a cumulative plot similar to those used in DNA sequence representation or analysis (Li 1997; Grigoriev 1998), for 22 human autosomes and 34 arms of metacentric chromosomes (Appendix Figure A1). The chromosome-scale or chromosome-arm scale recombination rate may be defined as the slope of a straight line that links the first and the last marker. For smaller chromosomes or chromosome arms, the end points in the Marey map tend to lie above the line with a slope equal to 1 (cM=Mb), i.e., smaller chromosomes(-arms) have larger cM/Mb ratios (see also Figure 16 of (Lander et al. 2001) and Table 12 of (Venter et al. 2001)).

We next regressed the genetic map length of chromosomes over their physical map length
Figure 1: Two-parameter regression of human genetic length over physical length. (A) Analysis at the chromosome scale. Female (red), male (blue), and sex-averaged (solid circle) genetic length of each chromosome (in cM) is plotted against its physical length (in Mb). The least-square regression lines are: \( y = 54.2 + 1.02x \) (female), \( y = 42.0 + 0.52x \) (male), \( y = 48.1 + 0.78x \) (sex-average). (B) Analysis of metacentric chromosome at the chromosome-arm scale. The best fit regression lines are: \( y = 29.0 + 1.05x \) (female), \( y = 27.1 + 0.48x \) (male), \( y = 28.0 + 0.77x \) (sex-average).

(Figure 1A), similar plot can be found in (Housworth and Stahl 2003). When sex-averaged, female and male genetic lengths are fitted separately, the three regression lines are described by:

\[
\begin{align*}
G_{ch, sex-ave, human} &= 48.1 + 0.78P \\
G_{ch, female, human} &= 54.2 + 1.02P \\
G_{ch, male, human} &= 42.0 + 0.53P
\end{align*}
\]

(1)

The normality assumption of regression residuals was tested graphically by a QQ-plot (Appendix Figure A2), and the normality condition does not seem to be violated.
Equation (1) shows that the \( y \)-intercept \( G_0 \) for female data is 29\% larger than for male data, whereas the slope \( k \) is 92\% larger. Thus, the different length of the male and female map mainly manifests as a different slope and less so as a different \( y \)-intercept. As can be seen from Figure 1(A), all human chromosomes exceed the minimal length of 50cM both for the male and the female genetic map.

To test the robustness of the \( y \)-intercept value, we added random noise to the genetic map length and repeated the regression analysis. The histogram of 50000 \( y \)-intercepts from this procedure is shown in Appendix Figure A3. Although values of \( G_0 \) range from 35 to 60, they are all far from zero.

We next repeated the analysis using chromosome arms instead of full chromosome as separate data points (Figure 1(B)). This leads to the regression equations:

\[
\begin{align*}
G_{\text{arm,sex-ave,human}} &= 28.0 + 0.77P \\
G_{\text{arm,female,human}} &= 29.0 + 1.05P \\
G_{\text{arm,male,human}} &= 27.1 + 0.48P. \\
\end{align*}
\]  

(2)

The \( y \) intercepts at chromosome arm-scale regression is now reduced to somewhat more than half of the intercept at the full chromosome scale. This reduction shows that cytogenetic constraints exert a smaller influence on the chromosome-arm scale than on the full chromosome scale.

Several methods can be used to show that the two-parameter regression model fits the data better than the one-parameter regressions. To this end, we first compared the coefficient of determination \( R^2 \) which is the proportion of variability explained by the regression model. The observed \( R^2 \) values of the one-parameter regression range between 0.48 and 0.87, whereas the \( R^2 \) values of the two-parameter regressions range between 0.86 and 0.98 (Table 1), indicating that the two-parameter regression explains more of the variability in the data.

We further cast the comparison between the one- and two-parameter regression as a model selection problem. Two such model comparison strategies are provided by the Akaike information criterion (AIC) (Akaike 1974) and Bayesian information criterion (BIC) (Schwarz 1978). Both AIC and BIC values for the two-parameter regression model are smaller than those for the one-parameter model, indicating a better statistical model (Appendix Table A1).
Finally, we tested the null hypothesis that $G_0$ is zero. The $p$-values in this test range between $10^{-13}$ and $10^{-8}$ (Table 1). Because the null hypothesis is that all chromosomes have the same RR, a simulated distribution of $G_0$ can be obtained from the regression over data that are obtained by the permuting of chromosome-specific cM/Mb ratios, while leaving the physical chromosome length unchanged. Out of 50000 such permutations, only two showed a $G_0$ value that is larger than the observed value of 48.1 (for sex-averaged full chromosome data), corresponding to a $p$-value of $4 \times 10^{-5}$. To summarize, all evaluation methods support the conclusion that the two-parameter regression model is better than the one-parameter model.

|                          | two-parameter | one-parameter | $p$-value for testing $G_0 = 0$ |
|--------------------------|---------------|---------------|-------------------------------|
| human chromosome, sex-averaged | $R^2$ = 0.976 | $R^2$ = 0.817 | $3.1 \times 10^{-10}$ |
| female                   | 0.984         | 0.866         | $8.6 \times 10^{-11}$         |
| male                     | 0.942         | 0.694         | $1.2 \times 10^{-8}$ |
| human chromosome arm, sex-averaged | $R^2$ = 0.955 | $R^2$ = 0.775 | $9.3 \times 10^{-13}$ |
| female                   | 0.964         | 0.861         | $7.1 \times 10^{-11}$         |
| male                     | 0.864         | 0.480         | $7.7 \times 10^{-11}$         |

Table 1: Comparison of the two-parameter and one-parameter regression models for human genetic length, at the chromosome scale (22 data points) and the chromosome-arm scale (34 data points): coefficient of determination ($R^2$) for 2- and 1-parameter regressions, and $p$-value for testing the null hypothesis of zero $y$-intercept.

As the deCode data were published more than six years ago, we further tested the chromosome-scale regression strategy on a more recent dataset, the Rutgers Map v.2 \cite{Matise2007}. The regression lines are $G = 53.33 + 0.87P$ (sex-average), $G = 50.29 + 1.19P$ (female), and $G = 57.58 + 0.57P$ (male), respectively. These results are consistent with the parameter estimations in Eq. (1), again showing that male and female data differ more in the slope than in the $y$-intercept.

Two other quantities can be derived from $G_0$ that help to interpret the $y$-intercept parameter. The first is the physical length $P_{min}$ on the regression line that corresponds to a specified minimum genetic length $G_{min}$ such that $G_{min} = G_0 + kP_{min}$. If we set $G_{min} = 50cM$, then we obtain $P_{min} = 2.45Mb$ for sex-averaged chromosome data. One may assume that for any
hypothetical chromosome with $P < P_{\text{min}}$, its genetic length $G$ remains constant at 50cM and does not decrease for shorter chromosome length. As the second quantity of interest, we define the percentage of genetic length that is explained by the inclusion of $G_0$ into the model as: 
$$\alpha = \frac{22G_0}{22G_0 + k \sum_{i=1}^{22} P_i}.$$  
For the sex-averaged data, we find that $\alpha = 31\%$ of variability is explained by the $y$-intercept. This value can also be obtained from the decomposition of 
$$RR = \frac{\sum_{i=1}^{22} G_i}{\sum_{i=1}^{22} P_i} = 22 \cdot \frac{G_0}{\sum_{i=1}^{22} P_i} + k: 1.13 = 0.35 + 0.78,$$  
because $0.35/1.13 = 31\%$. The relatively large percentage value once again highlights the importance of the $y$-intercept $G_0$ for modeling chromosome-scale recombination rate in human.

**Different intercept but similar slope in the two-parameter regression models for rat and mouse chromosomes**

Both rat (*Rattus norvegicus*) and mouse (*Mus musculus*) genome are known to have lower recombination rates than human ([Jense-Seaman et al. 2004](#)), with rat having a higher overall $RR$ than mouse. The rat genome has a roughly equal physical map length, but contains one more chromosome (n=20) than the mouse genome (n=19). Furthermore, rat chromosomes show a greater heterogeneity in their physical length and one may hypothesize that these karyotype differences contribute to the somewhat higher $RR$ in rat (0.62 cM/Mb vs. 0.57 cM/Mb in mouse). The regression models of the sex-averaged genetic length of rat and mouse chromosomes over their sequence lengths (Figure 2(A)) are:

$$G_{ch, sex-ave, rat} = 22.49 + 0.43P$$
$$G_{ch, sex-ave, mouse} = 15.62 + 0.44P.$$  
(3)

These models display a similar slope and the different overall $RR$ of rat and mouse mainly manifests as a different intercept value $G_0$.

Testing $G_0 = 0$ for the rat genome is significant ($p$-value= 0.0012), whereas testing $G_0 = 0$ for mouse genome (the fitted $G_0$ value for mouse is 69% of that for rat) is not significant ($p$-value = 0.11). AIC/BIC calculation confirms that the two-parameter regression is a convincingly better model for rat than the one-parameter regression, whereas this barely holds for the mouse data (Appendix Table A1). Thus, the mouse genome displays a non-significant excess of recombination on smaller chromosomes, which is consistent with the smaller variation of
chromosome size in the mouse genome. The greater $y$-intercept for the rat genome supports the hypothesis that cytogenetic factors contribute more to the genetic map length of rat than mouse.

Because the rat karyotype consists of both meta- and acrocentric chromosomes, we repeated the analysis after splitting all metacentric rat chromosome into two parts, based on the location of the centromere. Different from the human genome, for the rat genome this mainly affects the smaller chromosomes, which are often metacentric. The regression line is now described by $G = 7.48 + 0.52P$ and testing the intercept is still significant ($p$-value=0.024), though at a less stringent level. Thus, at the scale of chromosome-arms, the likelihood of crossovers in an interval is more determined by its physical length and less by influenced by any obligate recombination requirements.

**Recombination rate of small and large chromosomes in the chicken genome**

The chicken (*Gallus gallus*) genome consists of both large (macro-) and small (micro-) chromosomes (Hillier et al. 2004; Smith et al. 2000), with length ranging from a few Mb to close to 200Mb. The two-parameter regression model for the chicken genetic data in Figure 2(B) leads to:

$$G_{ch,sex-ave,chicken} = 34.68 + 2.79P.$$  \hspace{1cm} (4)

In this regression model, the non-zero intercept is significant with a $p$-value of $1.33 \times 10^{-7}$ and there is a considerable difference of AIC/BIC for the one- and two-parameter regression favoring the two-parameter model (Appendix Table A1). Both coefficients of determination for the 2- and 1-parameter regressions attain a high value: 0.98 and 0.93 respectively. A reason that the 1-parameter regression only marginally reduces the $R^2$ value is given by the fact that larger chromosomes contribute much more to the total variance, which is equally well captured by the 1-parameter model. Thus, two of the three methods confirm a relative excess of recombination on short chromosomes.

However, the orders of magnitude difference between the size of chicken chromosomes raises the question of the robustness of the regression. From (Hillier et al. 2004) and Appendix Figure A4, it is clear that the genetic length reaches a plateau at the level of 50cM for microchro-
somes smaller than 8Mb. When chromosomes below a certain length threshold are discarded from the regression analysis, the $y$-intercept value changes slightly, but not dramatically. For example, if the length thresholds for removal are 8Mb and 25Mb, $G_0$ for Eq. (4) decreases to 32.95 and 31.88. When the regression model is only fitted to the five largest chromosomes (longer than 50Mb), the model parameters are $G = 26.22 + 2.84P$. On the other hand, if we remove the largest five chromosomes, the regression line is $G = 31.86 + 3.01P$.

To see how the quality of the map distance measurements may influence these results, we next looked at the recently updated chicken map (Groenen et al. 2009), which contains more genetic markers and higher marker density. Applying the two-parameter regression leads to

$$G_{ch,sex-ave,chicken} = 34.23 + 2.04P.$$  \hfill (5)

As can be seen, the overall reduction of $RR$ as compared to the older map (Hillier et al. 2004) mainly manifests as reduced estimate of $k$, whereas the estimate of $G_0$ remains almost unchanged.

**Exceptionally high recombination on the largest honey bee chromosome leads to a better fit of the one-parameter than the two-parameter model**

Notably, the two-parameter regression does not provide a better fit for the genetic map data from honey-bee (*Apis mellifera*) (Beye et al. 2006) than the one-parameter model. When plotting the genetic length over physical length (Figure 2(C)), the $y$-intercept of the regression line does not significantly differ from zero ($p$-value = 0.81):

$$G_{ch,sex-ave,bee} = -4.22 + 23.49P.$$  \hfill (6)

The coefficient of determination for both the two- and one-parameter regression is around 0.95. In contrast to other genomes, AIC/BIC analysis favors the one-parameter regression model (Appendix Table A1).

As can be seen from Figure 2(C), the longest chromosome (chromosome 1) is four times the length of the shortest chromosome, and the regression result may depend on the presence of this “outlier”. To check this possibility, we repeated the analysis after chromosome 1 was removed, which led to the regression equation: $G = 28.71 + 20.36P$. However, also in this model,
testing $G_0 = 0$ is not significant ($p$-value=0.37), both the two- and one-parameter regressions exhibit similar coefficient of determination ($R^2 = 0.80, 0.79$), and the zero-intercept regression is still the better model according to AIC/BIC analysis (Appendix Table A1). Therefore, different from other species, the honey-bee genome does not display any significant excess of recombination on smaller chromosomes.

**Two-parameter regression at much shorter length scales: the example of budding yeast**

Yeast (\textit{S. cerevisiae}) has been extensively used to study the molecular machinery of recombination and it has a much smaller ($\sim 12$Mb) and more compact genome \cite{Cherry1997}. Although the physical length of yeast chromosomes only ranges from 200kb to 1.5Mb, their genetic length is between 100 and 500cM, even longer than the genetic length of human chromosomes. The best fitting regression line for the yeast genetic map is (Figure 2(D)):

$$G_{ch,sex-ave,yeast} = 49.12 + 284.74P.$$  \quad (7)

The non-zero $y$-intercept is significant ($p$-value = 0.009). The two-parameter regression is superior to the one-parameter model as judged by AIC/BIC (Appendix Table A1). The value of $y$-intercept, 49.12 cM is very close to 50cM which corresponds to almost one crossing over event for a hypothetical chromosome of physical length of zero.

The extremely high recombination rate in the yeast genome is surprising. From the molecular perspective, one can speculate about various hypotheses, such as a different meiotic regulatory system which makes a denser spatial distribution of chiasmata possible, a lack of secondary chromatin structure as compared to higher organisms so that the actual physical distance between two locations on chromosome is more or less equal to the linear sequence distance, or the lack of other supporting mechanism to hold chromatids together so that more chiasmata per chromosome arm are required for proper chromosome segregation. On the other hand, the $y$-intercept of the regression has a similar magnitude as that observed for higher organisms, indicating a similar relative excess of recombination on smaller chromosomes.
The difference between the central gene cluster and telomeric regions in worm genome is due to a difference in $G_0$

Finally, we used genetic map data from the worm *C. elegans* to show that the two-parameter regression strategy can also be useful to compare different regions within a genome. The chromosomes of *C. elegans* are unusual, because discrete centromeres are missing and the chromosomes are holocentric, i.e. microtubules attach at many sites for chromatid segregation (Tyler-Smith and Floridia [2000]). Accordingly, the Marey map analysis of the worm genome indicates that each worm chromosome can be partitioned in three regions: the central gene-rich region with a low recombination rate and two distal telomeric regions with high recombination rates (Barnes et al. [1995]). Therefore, we separately performed the regression analysis of genetic length over physical length for these two types of regions (Figure 3). The fitted regression coefficients are:

$$
G_{central,\text{worm}} = -2.22 + 1.01P
$$
$$
G_{distal,\text{worm}} = 18.39 + 0.94P
$$

(8)

Within the single parameter framework without the intercept term $G_0$, the two types of regions would have a very different cM/Mb ratio: 4.57 for telomeric regions, 0.68 for central regions. However, when allowing non-zero $G_0$ value, the two regions display similar slope values, 1.01 and 0.94. This indicates a constant excess of recombination in the distal region as compared to the central region in *C. elegans*, which is combined with a similar incremental cM/Mb ratio. Thus, after accounting for a fixed amount of recombination in a distal chromosome region, the likelihood of any additional recombination depends in similar strength on physical length in distal regions and central regions.

As a note of caution, one may point out that the regression coefficient in Eq. (8) is obtained from only a few data points. Nevertheless, further regression diagnostics supports our conclusion. For example, testing $G_0 = 0$ is significant for the distal regions ($p$-value= 0.006), but not significant for central regions ($p$-value=0.57). AIC/BIC analyses lead to the same conclusion (Appendix Table A1).
Discussion and Conclusions

Our results show that instead of the simpler genetic-to-physical length ratio, the relationship between the physical and genetic map length at chromosome scale is better described by a statistical model that contains a second parameter $G_0$, which is the $y$-intercept of the regression of genetic map length over the physical chromosome length. A conceptually similar approach was used earlier in measuring the genome-wide recombination rate of a species by counting the chiasmata on each chromosome in excess of one (Burt and Bell 1987).

The consideration of this intercept parameter is important, because karyotype structure has been established as an important determinant of genome-wide RR (De Villena and Sapienza 2001a; Coop 2005) and smaller chromosomes display higher RR (Lander et al. 2001; Hillier et al. 2004). Our proposed two-parameter model provides a formal expression of this size dependency of RR: $RR = G/P = k + G_0/P$, i.e., a constant term $k$ plus a second term that increases for smaller chromosome sizes $P$ (if $G_0$ is positive). This is what we observe for human, mouse, rat, chicken and yeast genomes. When writing $G_0$ as $G_0 = G - kP$, the $y$-intercept measures the amount of recombination after the physical map length has been accounted for. Therefore, one would expect that the total map length $G$ of a chromosome increases by $G_0$ after splitting it up into two separate parts. In fact, this has already been quantitatively observed for the experimental alteration of yeast chromosome I (Kaback et al. 1992).

When comparing RR between species, the usage of $k$ instead of the genome-wide cM/Mb ratio will reduce the influence of karyotype differences on the result. This was also the intention behind the counting of chiasmata per chromosome in excess of one (Burt and Bell 1987). In our study, the order of species remains unchanged, whether ranked by $k$ or by cM/Mb ratio. However, due to the different values of $k$, we cannot use a single regression line to model the genetic-physical length relationship across species. Thus, a molecular mechanism must exist that drives, within a particular species, the proportional increase of genetic over physical map length. This mechanism might typically act with weaker strength in larger genomes, which could contribute to the inverse correlation between genome size and RR (Lynch 2006).

Among mammals it was furthermore found that $RR$ is more similar for more closely related species (Dumont and Payseur 2008), which could be partly due to their similar karyotype.
It might be interesting to test where in the phylogenetic tree the signal might be altered, when using $k$ instead of the genome-wide cM/Mb ratio. In this context, it is also important that genome-wide $RR$ typically differs between genders and individuals (Broman et al. 1998; Kong et al. 2002, 2004; Cheung et al. 2007; Petkov et al. 2007). The biological factors that were invoked as possible explanations, such as differences in synaptonemal complex formation or crossover interference, may be more plastic than karyotype structure. The respective strength of these factors could also contribute to species differences and may be better measured by using $k$ than by using the genome-wide cM/Mb ratio.

If $k$ would equal to zero with the obligate chiasma requirement holding true, then $G_0$ were required to be 50cM. This pattern can be observed for female Opossum (Monodelphis domestica), where each chromosome acquires exactly one crossover near one of its telomeres (see Mikkelsen et al. 2007 and Appendix Figure A5). Similarly, very small chromosomes may always acquire exactly one crossover, despite reduced chromosome size, as seen for the microchromosomes in the chicken genome. In order to predict the transition from this plateau to the linear regression, we derived the minimum physical length parameter $P_{\text{min}}$ from a given $G_{\text{min}}$ and the estimated regression parameters. Note that if both physical and genetic length are measured as those in excess of $P_{\text{min}}$ and $G_{\text{min}}$, their ratio is exactly equal to $k$: 

$$\frac{(G-G_{\text{min}})}{(P-P_{\text{min}})} = \frac{(G-G_{\text{min}})}{(P-(G_{\text{min}}-G_0)/k)} = \frac{(G-G_{\text{min}})}{((kP+G_0-G_{\text{min}})/k)} = k.$$ 

Because reduced recombination may result in aneuploidy of smaller chromosomes (Warren et al. 1987; Brown et al. 2000), it is conceivable that the length of smaller chromosomes could influence genome-wide $RR$ by introducing a lower bound for the propensity for chiasma formation in a species. Our analysis supports the size of the smaller chromosomes as a strong determinant of genome-wide $RR$ for the six genomes studied in this paper (Appendix Figure A6). In log-log scale, the correlation coefficient between $RR$ and the shortest chromosome length is $-0.92$ ($p$-value= 0.008). If the recombination rate is measured by $k$, in log-log scale the correlation coefficient is $-0.91$ ($p$-value=0.01). This correlation is nearly as strong as the reported correlation between $RR$ and the total physical length for over 100 genomes ($cc= -0.99$, $p$-value= 0.0003 on log-log scale) as reported in (Lynch 2006). Obviously, data on more species are needed for a more conclusive analysis. Nevertheless, it may be interesting to point out that
the genome with the lowest known recombination rate, Opossum, lacks any short chromosome (Mikkelsen et al. 2007; Samollow et al. 2007).

Obviously, any genome-wide analysis relies on the availability of high quality data. We are convinced that the data used in this study are of sufficient quality to study recombination on the chromosomal scale. However, some error might be introduced by the fact that the used genetic maps are not perfect and, in particular for telomeres, missing some data. This can be seen for the chicken genomes, where the two chromosomes fall below the minimum genetic length of 50cM in the older map and climb to about 50cM in the newer map (Appendix Figure A4). Data selectively missing crossovers at the telomeres might lead to an underestimation of $G_0$ in the regression model.

We note that we restricted our analysis to chromosomes or chromosome-arms. If the genetic length is regressed over the length of much smaller regions, the coefficient of determination $R^2$ is expected to be much lower due to a mixture of recombination hot- and cold- spots. From a biological perspective, we also would not expect a positive $G_0$ value in such a regression, because there is no requirement for a Mb-sized region to have at least one chiasma to maintain meiotic integrity.

In summary, we find that the introduction of the $G_0$ parameter helps to understand the recombination rate differences between species, because it separates the effect of the requirement for at least one chiasma formation on smaller chromosomes from the factors that determine the amount of recombination on larger chromosomes. More specifically, the partitioning of chromosome-scale recombination rate leads to the following list of conclusions: i) human male-female $RR$ differences disproportionally affect larger chromosomes; ii) the higher recombination rate in the rat genome as compared to the mouse genome is likely to be caused by the higher number of smaller chromosomes that constitute the rat karyotype; iii) both chicken micro- and macro-chromosomes display a high $RR$ and the extraordinarily high $RR$ of some micro-chromosomes does not lead to an extraordinary excess of recombination on smaller chromosomes; iv) the honey-bee genome does not display any significant excess of recombination on smaller chromosomes; v) yeast displays a relative excess of recombination on smaller chromosomes that is similar to higher organisms, despite its outstandingly high overall recombination rate; vi) recombination of the worm genome mainly occurs in telomeric regions and given one
recombination per chromosome arm, the likelihood of a second recombination is determined by physical map length. These examples demonstrate that the proposed statistical framework allows to pinpoint differences of genomic recombination rate, which should be useful for the further study of genome-wide recombination rate as a quantitative trait of fundamental importance.

**Materials and methods**

**Genetic map data**

The human genetic map was obtained from (Kong et al. 2002) that uses 5136 microsatellite markers with 1257 meiotic events, and is estimated from pedigree data (Supplementary Table E: [http://www.nature.com/ng/journal/v31/n3/suppinfo/ng917_S1.html](http://www.nature.com/ng/journal/v31/n3/suppinfo/ng917_S1.html)). The rat (*Rattus norvegicus*) and mouse (*Mus musculus*) genetic map data were obtained from Table 1 of (Jense-Seaman et al. 2004), based on 2305 markers in rat and 4880 markers in mouse. The two chicken (*Gallus gallus*) genetic maps were obtained from Supplementary Table S2 of (Hillier et al. 2004), which is built from 1471 markers, and from Table 1 of (Groenen et al. 2009) built from 9258 markers. The honey bee (*Apis mellifera*) genetic map was obtained from Table 2 of (Beve et al. 2006) based on 1500 markers. The budding yeast (*Saccharomyces cerevisiae*) genetic map was downloaded from [http://downloads.yeastgenome.org/chromosomal_feature/SGD_features.tab](http://downloads.yeastgenome.org/chromosomal_feature/SGD_features.tab). The worm (*Caenorhabditis elegans*) physical and genetic lengths of central “gene clusters” and distal “arms” were obtained from Table 1 of (Barnes et al. 1995) based on 168 markers.

**Measuring how good a linear regression is by coefficient of determination**

Regression analyses were carried out by the *lm*() subroutine in *R* statistical package. For genetic lengths, \( \{G_i\} \) \((i = 1, 2, \ldots n)\), e.g., \(n = 22\) for the chromosome-scale regression and \(n = 34\) for the chromosome-arm-scale regression), one can regress them over sequence lengths
\{P_i\} (i = 1, 2, \cdots n) allowing y-intercept (non-zero G when P approaches 0):

\[ G = G_0 + kP, \]

(9)

or, without the y-intercept (G approach 0 as P approaches 0):

\[ G = kP. \]

(10)

How good a linear regression model fits the data can be measured by the coefficient of determination \(R^2\), which is the proportion of variability that is explained by the model. More specifically, if \(SS_{tot} = \sum_{i=1}^{n}(G_i - E[G_i])^2\) is the total sum of squares of the genetic lengths of chromosomes, the term \(SS_{err} = \sum_{i=1}^{n}(G_i - G_0 - kP_i)^2\) for allowing non-zero y-intercept, or the term, \(SS_{err} = \sum_{i=1}^{n}(G_i - kP_i)^2\) for not allowing y-intercept, is the residual sum of squares (RSS), then

\[ R^2 = 1 - \frac{SS_{err}}{SS_{tot}} = 1 - \frac{RSS}{SS_{tot}} \]

(11)

Model selection by Akaike information criterion

Akaike information criterion (AIC) of a statistical model is defined as \(2p - 2\log(L)\) where \(p\) is the number of parameters in the model, and \(L\) is the maximum likelihood estimated from the data. Similarly, Bayesian information criterion (BIC) is defined as \(\log(n)p - 2\log(L)\), where \(n\) is the number of samples used to calculate the likelihood. For linear regression, AIC/BIC is related to the residual sum of squares (RSS) according to (Venables and Ripley 1999) by:

\[ \text{AIC} = 2p + n\log(RSS/n) \]

\[ \text{BIC} = \log(n)p + n\log(RSS/n) \]

(12)

where \(n\) is the number of sample points for the regression analysis. Between two statistical models that are fitted to the same dataset, the model with a smaller AIC/BIC value is considered to be better than the model with a larger AIC/BIC value.

For the comparison between the two- and one-parameter regressions, we have:

\[ \text{AIC}_2 - \text{AIC}_1 = 2 - n\log \frac{RSS_1}{RSS_2} \]

\[ \text{BIC}_2 - \text{BIC}_1 = \log(n) - n\log \frac{RSS_1}{RSS_2} \]

(13)
If the second term, \( n \log(RSS_1/RSS_2) \), is larger than two (for AIC, or \( \log(n) \) for BIC), then the two-parameter regression can be seen as the better model than the single-parameter regression.

Quantities derived from \( G_0 \)

The linear relationship between \( G \) and \( P \) cannot extend to the physical length of zero, if the \( y \)-intercept is greater than zero and the obligate chiasma requirement holds. Therefore, a point \( P_{min} \) must exist below which genetic map length remains constant at \( G_{min} \), independent from the actual physical map length of a chromosome. We can define this transition point as follows: \( P_{min} \) is the physical length for which the regression line crosses the horizontal line defined by the minimum genetic length \( G_{min} \), thus \( P_{min} = (G_{min} - G_0)/k \).

Another derived quantity is the genome-wide percentage of genetic length that is explained by \( G_0 \). For a single chromosome \( (i) \), this percentage is \( \alpha_i \equiv G_0/(G_0 + kP_i) \). For the whole genome, it is \( \alpha \equiv nG_0/(nG_0 + k \sum_i P_i) \), where \( n \) is the number of chromosomes. This definition of \( \alpha \) is valid only when the \( y \)-intercept is positive \( (G_0 > 0) \).

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Appendix

1 Marey Map for human chromosomes (Fig. A1)
2 Checking the normality assumption of the regression (Fig.A2)
3 Testing the robustness of regression result by adding random noise to the genetic map (Fig.A3)
## AIC and BIC of two-parameter regression models vs. one-parameter models (Table A1)

| source of genetic length data                      | $\Delta AIC = AIC_2 - AIC_1$ | $\Delta BIC = BIC_2 - BIC_1$ |
|---------------------------------------------------|-------------------------------|-------------------------------|
| human chromosome, sex-averaged                    | -42.5                         | -41.4                         |
| female                                           | -45.3                         | -44.2                         |
| male                                             | -34.5                         | -33.4                         |
| human ch. arm, sex-averaged                       | -52.9                         | -51.4                         |
| female                                           | -43.8                         | -42.3                         |
| male                                             | -43.6                         | -42.1                         |
| rat, chromosome, sex-averaged                     | -9.9                          | -8.9                          |
| mouse, chromosome, sex-averaged                   | -1.1                          | -0.2                          |
| chicken, chromosome, sex-averaged                 | -28.7                         | -27.5                         |
| honeybee, chromosome, sex-averaged                | 1.9                           | 2.7                           |
| remove chromosome 1                              | 1.0                           | 1.7                           |
| yeast, chromosome, sex-averaged                   | -6.1                          | -5.3                          |
| worm, central region                              | 1.4                           | 0.96                          |
| distal region                                     | -7.9                          | -7.6                          |

Table A1: Difference of Akaike and Bayesian information criterion (AIC and BIC) between the two- and one-parameter regression models for modeling chromosome-scale recombination in 7 genomes. A negative $\Delta AIC$ or $\Delta BIC$ value favors the two-parameter regression model.
5 Chicken genetic length vs. physical length in log-log scale (Fig.A4)
6 Two-parameter regression model of Opossum chromosome genetic length (Fig.A5)
7 Recombination rates of six genomes as a function of the smallest chromosome size (Fig. A6)
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Figure 2: The genetic length (in cM) vs. physical length (in Mb) plotted for five genomes: (A) mouse (crosses) (*Mus musculus*) and rat (circles) (*Rattus norvegicus*). Source: Table 1 of (Jense-Seaman et al. 2004). The regression lines are: $y = 15.62 + 0.44x$ (mouse), $y = 22.49 + 0.43x$ (rat). (B) chicken (*Gallus gallus*). Source: old data (year 2004, circle) is from supplementary Table A2 of (Hillier et al. 2004); new data (year 2008, cross) is from Table 1 of (Groenen et al. 2009). The regression line is: $y = 34.68 + 2.79x$ (old data) and $y = 34.23 + 2.04x$ (new data). (C) honeybee (*Apis mellifera*). Source: Table 2 of (Beye et al. 2006). The regression line is $y = -4.22 + 23.49x$. (D) budding yeast (*Saccharomyces cerevisiae*). Source: [http://downloads.yeastgenome.org/chromosomal_feature/SGD_features.tab](http://downloads.yeastgenome.org/chromosomal_feature/SGD_features.tab) The regression line is $y = 49.12 + 287.74x$. 
Figure 3: The cM-Mb plot using the physical and genetic length of central gene clusters (5 data points) and distal arms (10 data points) of five worm (Caenorhabditis elegans) chromosomes (Table 1 of (Barnes et al. 1995)). The best fitting regression lines are $y = 18.39 + 0.94x$ for the distal/telomeric arms (crosses), and $y = -2.22 + 1.01x$ for the central gene cluster regions (circles).
Figure A1: Marey map for human genetic data (Supplementary Table E of (Kong et al., 2002), with y-axis showing the genetic distance (in cM) from the first marker to the current marker, and x-axis showing the physical distance (in Mb). (A) Each line traces a chromosome (chromosome name is shown as label). (B) Each line traces an arm of a meta-centric chromosome (p- and q-arms are shown as label). The straight line indicates cM=Mb.
Figure A2: QQ-plots of the residuals of regression ($\epsilon = G - G_0 - kP$) against simulated normal variables. The variance of the normal variables is chosen to be equal to that of the regression residuals. Due to the small number of sample points (22 for human chromosomes, 34 for human chromosome arms), four sets of normal random variables were generated to test the robustness. In each QQ-plot, “o” denotes the regression for the sex-averaged map of human chromosomes, “f” for the human female map data, “m” for human male map data, and “a” for sex-averaged human map data for chromosome arms (see Eqs.(1,2)).
Figure A3: Histogram of $G_0$ for human sex-averaged chromosome regression $G_{ch,sex-ave,human} = G_0 + kP$ when noise is added to the genetic length $G$. The noise was modeled as a normally distributed variable with zero-mean and standard deviation (sd) of 8.51517, which is the observed sd of regression residuals $\epsilon = G - G_0 - kP$, for $G_{ch,sex-ave,human} \sim P$ in Eq.(1).
Figure A4: The genetic length (in cM) vs. physical length (in Mb) plots for chicken genome (*Gallus gallus*) in log-log scale (for linear-linear scale, see Fig. 2(B)).
Figure A5: Two-parameter regression of Opossum *Monodelphis domestica* chromosome genetic length (cM) over physical length (Mb). The regression coefficients for female map (red) are: $G_{female} = 54.206 + 0.003P$, for male map (blue): $G_{male} = 19.610 + 0.216P$, and for sex-averaged map (black): $G_{sex-ave} = 31.275 + 0.114P$. 
Figure A6: Recombination rates of six genomes as a function of the smallest chromosome size. Genome-wide recombination rate is measured both by the genome-wide genetic-to-physical length ratio ($RR = G/P$) (circles) and by the regression slope $k$ (pluses). The plot is in log-log scale.