CONY: A Bayesian procedure for detecting copy number variations from sequencing read depths

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Copy number variations (CNVs) are genomic structural mutations consisting of abnormal numbers of fragment copies. Next-generation sequencing of read-depth signals mirrors these variants. Some tools used to predict CNVs by depth have been published, but most of these tools can be applied to only a specific data type due to modeling limitations. We develop a tool for copy number variation detection by a Bayesian procedure, i.e., CONY, that adopts a Bayesian hierarchical model and an efficient reversible-jump Markov chain Monte Carlo inference algorithm for whole genome sequencing of read-depth data. CONY can be applied not only to individual samples for estimating the absolute number of copies but also to case-control pairs for detecting patient-specific variations. We evaluate the performance of CONY and compare CONY with competing approaches through simulations and by using experimental data from the 1000 Genomes Project. CONY outperforms the other methods in terms of accuracy in both single-sample and paired-samples analyses. In addition, CONY performs well regardless of whether the data coverage is high or low. CONY is useful for detecting both absolute and relative CNVs from read-depth data sequences. The package is available at https://github.com/weiyuchung/CONY.

Copy number variations (CNVs) are genomic structural mutations consisting of abnormal numbers of deoxyribonucleic acid (DNA) section copies. CNVs were originally defined to range from one kilobasepair to several megabasepairs1-2 and widened to include small variants that are larger than 50 basepairs in size3,4. Currently, approximately 7 million CNVs identified in 1 million variant regions are catalogued in the Database of Genomic Variants (DGV)5,6. Half the identified CNVs overlap with protein-coding regions, which results in gene expression changes7. CNVs have been confirmed to play important roles in human diseases; for example, glycoporphin CNVs in malaria resistance8, beta-defensin CNVs in Psoriasis9,10, CNVs in 15q11.2 for the perigenual anterior cingulate cortex in schizophrenia and Alzheimer's disease11,12, and some pathogenic CNVs in developmental delay, autism spectrum disorders, and various congenital malformations13,14. Furthermore, somatic copy number aberrations have been considered to be associated with human cancers and to categorize the subtypes of cancer15, such as breast cancer16,17, lung cancer18,19, and colorectal cancer20,21.

Array comparative genomic hybridization22,23 and single nucleotide polymorphism arrays24,25 have been used to detect CNVs over the past few years; however, the boundaries of CNVs cannot be explicitly identified due to the sparse probe coverage. Recently, next-generation sequencing (NGS)26,27 has provided a more accurate option for CNV identification and breakpoint prediction through the high-resolution analysis of sequential DNA nucleotide bases. Various strategies, including read depth28-36, paired-end mapping37-40, split read41-43, assembly44-46 and integrative approaches47-51, have been adopted to detect CNVs in NGS data. Read depth analysis becomes a major method because of less restriction for read lengths and insert sizes26,27,52, which are critical limitations for other strategies. Besides, depth data can be derived from both paired- and single-end sequencing reads with appropriate mapping and normalizing procedures.

In the read-depth approach, CNV identification assumes that the number of reads is proportional to the number of DNA copies. Hypothesis testing, change point segmentation, and the hidden Markov model are commonly used methods in this field. While many practical tools have been developed using these types of statistical algorithms, the link between sequencing depth information and CNVs is not completely understood. In hypothesis testing methods, each depth is independently tested for a significant CNV35,36, but correlation of depths should

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be considered through the corresponding genomic locations. The adjustment methods used for multiple testing issues also need to be evaluated rigorously. In change point algorithms, copy number (CN) regions are first identified by a segmentation algorithm, and then the states of the proposed CN regions are estimated. However, the performance of the segmentation step has an obvious impact on the downstream CNV detection accuracy. To overcome these shortcomings, a statistical model approach that considers genetic information from whole genome sequencing depths to simultaneously identify CN regions and states is presented in this paper.

In addition, most existing approaches were proposed for a specific sample design. Single-sample analyses can estimate absolute CN calls, and are implemented in personalized medicine. However, read data from one single sample only contain individual genomic information, not population-level variations; as a result, it is not easy to find the potential biases especially in the low coverage data. In contrast, depth ratios of paired samples (case/control or tumor/normal) identify patient-specific relative CNVs and are conveniently utilized in association studies. While background or platform noises may be efficiently eliminated through the comparative depths, combining sample information from different sequencing coverages or platforms remain difficult issues. The proposed model-based algorithm in this study could be applied to various sample designs to fulfill the thorough data transformation and the parameter settings.

Given the aforementioned challenges, we propose a comprehensive approach called copy number variation detection via a Bayesian procedure (CONY). A Bayesian hierarchical model is constructed to integrate the sequencing depth signals, the corresponding genomic position, and the potential CNV information. The efficient sampling algorithm, i.e., reversible-jump Markov chain Monte Carlo (RJMCMC), is modified to infer the states and breakpoints of the CN regions. An appropriate analytic section length of the genome for the RJMCMC algorithm is suggested to reduce the unbalanced effects that result from the extreme difference between normal and variant region sizes. The usefulness of the CONY algorithm is demonstrated by both simulations and an analysis of experimental data from the 1000 Genomes Project.

Materials

The 1000 Genomes Project. Whole genome sequencing data of two samples NA12156 and NA12878 (SRA accessions ERX000125 and ERX000080, respectively) provided by the 1000 Genomes Project were analyzed. Each of the samples was used to identify the absolute CNVs, and they were matched to form case/control pairs (NA12156/NA12878 and NA12878/NA12156) to identify the case-specific relative CNVs. The identified CNVs were compared with CNV lists reported in the Database of Genomic Variants. Sequencing reads generated by the Illumina platform with 4.1 to 5.7X coverage and mapped to the human genome 19 (hg19/GRCh37) reference genome with default adjustments were downloaded from the 1000 Genomes Project ftp.

Another two experimental samples HG00419 and HG01595 from the project, which were sequenced with both low (5.2 to 9.8X as SRA accessions SRX724413 and SRX720422, respectively) and high (33.6 to 35.4X as SRA accessions SRX500074 and SRX50114, respectively) coverages, were also analyzed to show consistency of results from CONY across samples and evaluate the coverage effect. Both samples were used for the single-sample analysis; HG00419/HG01595 and HG01595/HG00419 were matched for the paired-samples analysis. Reads mapped to the hg38/GRCh38 human reference genomes were adopted.

The simulation study. In the single-sample analysis, DNA sequences were generated from one hundred samples with predetermined CNVs. We used the hg19 chromosome 20 (chr20) as the template. The template sequence was copied to one strand and deleted/duplicated in pieces to mimic the copy loss/gain to the other strand for each sample. Twenty pieces for copy losses were deleted from the variant strand as copy number (CN) 1, and twenty pieces for copy gains were randomly duplicated 1, 2, or 3 times as CN 3, 4, or 5, respectively. The artificial pieces were set at 10 different sizes (1, 2.5, 5, 10, 25, 50, 100, 250, 500, and 1000 kilobases (kb)) using 2 of each for the copy losses/gains. The synthetic CNV regions accounted for 12% of the human genome, which is consistent with a recent report. In the paired-samples analysis, simulated samples from the single-sample analysis were used as case samples. One common control sample sequence was copied from the hg19 chr20 template for both strands. In total, two million paired-end reads with a length of 70 bases (bp) and a coverage of 2.2X (low coverage) or 22X (high coverage) were generated for each sample via the sequencing simulation software Wgsim. The simulated reads were aligned to the reference genome by BWA and subjected to data preprocessing.

Methods

A Bayesian model-based procedure, i.e., CONY, that is able to identify both absolute and relative CNVs from both single-sample and paired-samples DNA sequencing is proposed. In this procedure, read-depth signals (RDSs) derived from preprocessed sequencing reads are used to estimate CNVs via a Bayesian hierarchical model and the RJMCMC algorithm.

The sequencing reads are aligned with the reference genome, subjected to preprocessing steps, and accumulated as read depths per base via published tools. Next, the base-read depths in a small contiguous region (referred to as a window) are summed as the window read depth of each sample. After adjusting for potential biases, the window read depths are transformed to RDSs by logarithm (single-sample analysis) or log-ratio (paired-samples analysis) equations. RDSs are linked to the states and breakpoints of CN regions via a comprehensive Bayesian hierarchical model. A modified RJMCMC algorithm is constructed to generate samples for parameter inferences with two main moves (updating CN states and updating boundaries) and four jumping strategies (merge, split, trifid, and boundary change) for updating the boundaries. After 5,000 burn-ins, the windows with the abnormal CNs are tested via Bayes factors in each additional 1,000 iterations until full coverage is achieved. The details of the CONY procedure are provided in the following discussion, and a flow chart of the analysis is depicted in Fig. 1.
Read alignment, data preprocessing, and read-depth signal calculation. First, the decoded sample sequencing reads (FASTQ format) are aligned with the reference sequence (FASTA file) to ensure the corresponding locations in the genome via commonly used tools, such as BWA and Bowtie2. The best-matched position information of each read is written in SAM/BAM format using SAMtools software. The low-quality reads and experimental duplicates are removed, including base-calling quality scores lower than 13, mapping quality scores lower than 30, and PCR duplicates. Then, the good-quality reads are piled to obtain accumulated measurements of each nucleotide, which are referred to as the “base-read depth.”

Base-read depths are insufficient for identifying CNVs with high specificity. The potential systematic biases easily override the true CNV evidence because of the weak information from a single base. Moreover, a single signal has insufficient statistical strength to support the assumption of a uniform relationship between the CNVs and read depth. To increase the power of the read-depth information, the summarized signals from several bases are considered. A series of consecutive bases constitute a window, and the depths of the bases within the window are accumulated to obtain stable and convincing read-depth information.

The genome is partitioned into nonoverlapping sliding windows with an equal size of 100 bp as a default, and the base-read depths in each window are summed as the raw “window read depth.” The raw window read depth is denoted by \( R_{Raw,i} \).

Two major bias effects (i.e., the percentage of indefinable bases and GC content) should be adjusted to strengthen the evidence of CNVs in the raw window read depths. First, the percentage of bases with N code (i.e., indefinable bases) should be considered. Because no depths are counted for these indefinable bases, the window read depths should be adjusted to balance the information across windows. Then, the \( i \)th window read depth is adjusted by the following equation:

\[
R_{CorrSize,i} = R_{Raw,i} \times \frac{\text{window size}}{\text{number of indefinable bases in the } i \text{th window}}.
\]

Second, the GC content is a notable source of noise in the depth estimation, especially using the Illumina platform. The method used for the GC content adjustment follows that of a published study. The GC-adjusted window read depth is calculated by the following formula if the percentage of G and C codes in the \( i \)th window is in the range from 20% to 80%:

\[
R_{CorrGC,i} = R_{CorrSize,i} \times \frac{R_{GC,i}}{R_{GC,avg}},
\]

where \( R_{GC,i} \) and \( R_{GC,avg} \) represent the predicted depth in the \( i \)th window via a local regression (LOESS) and the average depth over all windows. Regarding the LOESS model settings, the proportion of neighborhood points is spanned to 75%, and the weight follows a typical tri-cubic function. Since the LOESS adjustment does not work for extreme

Figure 1. Flowchart of read alignment, data preprocessing and CNV detection.
GC percentages (<20% or >80%), the depths of these windows are not adjusted; thus, \( R_{\text{corrGC},i} = R_{\text{corrSize},i} \). Furthermore, windows with more than half indefinable codes or zero depth are marked and excluded from further analysis. Then, the window read-depth signals (hereafter referred to as RDSs, \( D_i \)) for the single-sample and paired-samples analyses are calculated by logarithm (i.e., \( D_i = \log(R_{\text{corrGC},i}) \)) and log-ratio (i.e., \( D_i = \log(R_{\text{corrGC},i}(\text{Case})/R_{\text{corrGC},i}(\text{Control})) \)) transformations.

**Bayesian hierarchical model.** Following the process outlined above, the adjusted window RDSs were prepared for a downtrend application to estimate the CNVs. A Bayesian hierarchical model is proposed for detecting the absolute/relative CNVs from single-sample/paired-samples window RDSs.
This model aims to divide the whole genome with I windows into M CN regions to group consecutive windows with the same underlying CN. The comprehensive structure constructs the relationships among the window RDSs \( \mathbf{D} = \{ D_1, D_2, \ldots, D_I \} \), CN states underlying CN regions \( \mathbf{C} = \{ C_1, C_2, \ldots, C_M \} \) and CN breakpoint indicators of window boundaries \( \mathbf{B} = \{ B_0, B_1, \ldots, B_I \} \) (Fig. 2). The Bayesian approach starts with the prior belief that the parameters follow the prior distribution \( p(\mathbf{B}, \mathbf{C}) \) and uses the likelihood from data \( p(\mathbf{D} | \mathbf{B}, \mathbf{C}) \) to update the parameters a posterior. Unlike some existing tools, our proposed Bayesian hierarchical model comprehensively considers parameter relations across the analytic genome rather than just information in consecutive windows. The inferences are based on the posterior distribution \( p(\mathbf{B}, \mathbf{C}, \mathbf{D}) \), which is proportional to the priors multiplying the data likelihood, \( p(\mathbf{B}, \mathbf{C}, \mathbf{D}) \propto p(\mathbf{B}, \mathbf{C}) \times p(\mathbf{D} | \mathbf{B}, \mathbf{C}) = p(\mathbf{B}) \times p(\mathbf{C} | \mathbf{B}) \times p(\mathbf{D} | \mathbf{B}, \mathbf{C}) \). Three parts are included in factorization, including the window boundary \( p(\mathbf{B}) \), CN state \( p(\mathbf{C} | \mathbf{B}) \), and depth \( p(\mathbf{D} | \mathbf{B}, \mathbf{C}) \). The details of factorization and the hyperparameter settings are shown below.

Window boundary part \( p(\mathbf{B}) \). Parameter \( \mathbf{B} = \{ B_0, B_1, \ldots, B_I \} \) is used to represent whether the window boundaries are the breakpoints of the CN regions. \( B_i = 1 \) if windows \( i \) and \( i+1 \) have different underlying CNs for \( i = 1, 2, \ldots, I-1 \) (i.e., the \( i \)th window boundary is the breakpoint of two CN regions). Otherwise, \( B_i \) is denoted by 0. \( B_0 \) and \( B_I \) are set to 1 due to the left and right borders. Assume that \( B_i \) follows an independent Bernoulli distribution with success probability \( \lambda \). The probability of the window boundaries is \( p(\mathbf{B}) = p(\mathbf{B}_0, \mathbf{B}_1, \ldots, \mathbf{B}_I) = \lambda^{M-1} \times (1 - \lambda)^{I-M}, \) where \( M = \sum_{i=0}^{I} B_i \). Thus, there is a quantity \( M \) of \( B_i \) with a value of 1 for \( i \) from 1 to \( I \), and the genome is separated into \( M \) CN regions.

Copy number state part \( p(\mathbf{C} | \mathbf{B}) \). The whole genome is divided into \( M \) CN regions when breakpoints \( \mathbf{B} \) are given. Next, the CN states of each region \( \mathbf{C} = \{ C_1, C_2, \ldots, C_M \} \) are described based on conditional probability \( p(\mathbf{C} | \mathbf{B}) = p(C_1, \ldots, C_M | \mathbf{B}) \), which can be factored as \( p(\mathbf{C}_1 | \mathbf{B}) \times p(\mathbf{C}_2, \mathbf{C}_3 | \mathbf{B}) \times \ldots \times p(\mathbf{C}_M | \mathbf{C}_1, \ldots, \mathbf{C}_{M-1}, \mathbf{B}) \). Because the consecutive CN regions must have different states, the state of each region is restricted to the adjacent sides. Therefore, the conditional probability is simplified as \( p(\mathbf{C}_1 | \mathbf{B}) \times p(\mathbf{C}_2, \mathbf{C}_3 | \mathbf{B}) \times \ldots \times p(\mathbf{C}_M | \mathbf{C}_1, \ldots, \mathbf{C}_{M-1}, \mathbf{B}) \).

For the state of the first region \( C_1 = \{ C_{1,0}, C_{1,1}, \ldots, C_{1,K} \} \), a one-trial multinomial distribution with a prespecified category number \( K \) is adopted, i.e., \( C_{1,i}, C_{1,1}, \ldots, C_{1,K} \sim \text{Multinomial}(1; \mathbf{W}_{1,0}, \mathbf{W}_{1,2}, \ldots, \mathbf{W}_{1,K}) \). The element \( C_{1,i} \) of \( \mathbf{C}_1 \) is equal to 1, then the CN state of the first region is denoted by \( C_{1,1} \). The weight vector \( \mathbf{W}_{1,0} = [\mathbf{W}_{1,0}, \mathbf{W}_{1,2}, \ldots, \mathbf{W}_{1,K}] \) follows a conjugate Dirichlet distribution with hyperparameter \( \mathbf{W}_{0} = [\mathbf{W}_{0,0}, \mathbf{W}_{0,2}, \ldots, \mathbf{W}_{0,K}] \).

The state of the other regions must be different from the previous state based on the above conditional probability factorization. Assuming the state of the \( (m-1) \)th region is \( k \) (i.e., \( C_{m-1,k} = 1 \) or \( C_{m-1,k} = k \)), the state of the \( m \)-th region \( C_m = \{ C_{m,1}, C_{m,2}, \ldots, C_{m,k-1}, C_{m,k+1}, \ldots, C_{m,C_K} \} \) could decrease by one dimension with \( K-1 \) categories. \( C_m \) follows a one-trial multinomial distribution with weight vector \( \mathbf{W}_m = [\mathbf{W}_{m,1}, \mathbf{W}_{m,2}, \ldots, \mathbf{W}_{m,k-1}, \mathbf{W}_{m,k+1}, \ldots, \mathbf{W}_{m,K}] \) and the weight is Dirichlet distributed with parameter \( \mathbf{W}_m = [\mathbf{W}_{0,1}, \mathbf{W}_{0,2}, \ldots, \mathbf{W}_{0,K}] \). The hyperparameter \( \mathbf{W}_0 = [\mathbf{W}_{0,1}, \mathbf{W}_{0,2}, \ldots, \mathbf{W}_{0,K}] \) of weight \( \mathbf{W} = [\mathbf{W}_{0,1}, \mathbf{W}_{0,2}, \ldots, \mathbf{W}_{0,K}] \) is estimated via the empirical method introduced in Supplementary Text 1 (Hyperparameters setting).

The conditional probability of the CN states given the breakpoints is summarized as follows:

\[
P(\mathbf{C} | \mathbf{B}) = \int p(\mathbf{C}_1 | \mathbf{B}, \mathbf{W}) \times p(\mathbf{C}_2, \mathbf{C}_3 | \mathbf{B}, \mathbf{W}) \times \cdots \times p(\mathbf{C}_{M-1}, \mathbf{C}_M | \mathbf{B}, \mathbf{W}) \times p(\mathbf{W} | \mathbf{B}) d\mathbf{W}
\]

where \( n_{k} \) is the number of regions located after the regions with CN state \( k \), and \( n_{k,k} \) is the number of regions with state \( k \) among these \( n_{k} \) regions. Based on this formula, this model connects information not only from these regions with identical CN states but also from the same previous regions to strengthen the state relationship.

In addition, the number of state categories \( K \) needs to be pre-assigned in this procedure. For a single-sample analysis, the states represent the absolute CN, and we set \( K = 5 \) as the default. For paired samples, the states represent the relative CN, and we set \( K = 3 \) as the default, representing copy loss, normal and copy gain statuses.

Depth part \( p(\mathbf{D} | \mathbf{B}, \mathbf{C}) \). Given the breakpoints and states of each CN region, we assume that RDSs within the same copy number region follow an independent normal distribution with a common mean and variance. Moreover, the normal and inverse-gamma conjugate priors connect windows from different CN regions that belong to the same CN state. Therefore, the conditional likelihood is derived as follows:
where $L_m$ is defined as the number of windows in CN region $m$, and $L_m = 0$. The settings of hyperparameters $\mu$, $\alpha$, $\beta$, and $\kappa$ are shown in Supplementary Text 1 (Hyperparameters setting).

**Proportional posterior distribution.** By multiplying the window boundary, CN state, and depth parts mentioned above, the proportional posterior distributions of $\mathbf{B}$ and $\mathbf{C}$ are obtained.

$$p(\mathbf{D} | \mathbf{B}, \mathbf{C}) = p(D_1 | \mathbf{C}, \mathbf{B}) \times p(D_2 | \mathbf{C}, \mathbf{B}) \times \cdots \times p(D_i | \mathbf{C}, \mathbf{B})$$

$$= \int \int \int \int \int p(D_1 | \mathbf{C}, \mathbf{B}, \mu_1, \sigma_1^2) p(\mu_1, \sigma_1^2) d\mu_1 d\sigma_1^2$$

$$\times \int \int \int \int p(D_2 | \mathbf{C}, \mathbf{B}, \mu_2, \sigma_2^2) p(\mu_2, \sigma_2^2) d\mu_2 d\sigma_2^2$$

$$\times \cdots$$

$$\times \int \int \int \int p(D_i | \mathbf{C}, \mathbf{B}, \mu_i, \sigma_i^2) p(\mu_i, \sigma_i^2) d\mu_i d\sigma_i^2$$

$$= \prod_{i=1}^{I} \left[ \int \int N(D_i | \mu_i, \sigma_i^2) \times N \left( \mu_i | \mu_{0i}, \sigma_{0i}^2/k_{C_i}^2 \right) \times IG(\sigma_i^2 | \alpha_C, \beta_C) d\mu_i d\sigma_i^2 \right]$$

$$= \prod_{m=1}^{M} \left[ \int \int \frac{(-1)^m}{\sqrt{C_m^2 + L_m}} \times \frac{\Gamma(\alpha_C + L_m/2)}{\Gamma(\alpha_C^2)} \frac{\beta_C^{\alpha_C}}{\beta_{C_m}^{\alpha_C + L_m}} \right.$$

$$\times \left. \left[ \frac{\kappa_{C_m}^2}{C_m^2} + \sum_{i=1}^{L_m-1} \frac{\kappa_{C_m}^2}{C_m^2} \right] \right]$$

The relationships among the parameters are depicted in Fig. S1.

**Reversible-jump Markov chain Monte Carlo algorithm.** Two groups of variables, i.e., CN states $\mathbf{C}$ and window boundaries $\mathbf{B}$, are estimated from the derived posterior distribution $p(\mathbf{B}, \mathbf{C} | \mathbf{D})$. In our model, the number of parameters is not fixed, primarily because the values of $\mathbf{B}$ can affect the numbers of CN regions and corresponding states $\mathbf{C}$. A powerful algorithm, i.e., RJMCMC$^{55}$, is adopted for sampling from a specified distribution with a variable number of dimensions. We construct a RJMCMC algorithm with two efficient moves, i.e., "Update copy number states $\mathbf{C}$" and "Update window boundaries $\mathbf{B}$" for each transition. The details are illustrated below.

To update CN states $\mathbf{C}$, all analyzed regions are updated together via a Gibbs sampler. Conditional on the values of boundaries $\mathbf{B}$ and RDDSs $\mathbf{D}$, the probabilities of all possible CN state combinations are calculated. The combination with the maximum probability is selected. The conditional probability is expressed as follows:
However, updating window boundaries $B$ is complex. Because the values of the window boundaries are subject to the dimension of the CN regions and corresponding states, not only the boundaries but also the neighboring CN states are updated in this move. To explore the parameter space efficiently and completely, four novel jumping strategies are adopted: merge, split, trifid, and boundary change. The relationships among the jumping strategies are illustrated in Fig. 3.

In the merge strategy, one window boundary with value 1 (i.e., CN region breakpoint) is randomly changed to a value of 0, and then, the adjacent CN regions sharing the selected boundary are combined. The state of the new CN region is chosen from two states of the original CN regions with equal probability. Assume that the $m$ and $m+1$ regions are merged into a new region with index $m^*$. Then, the candidate status is accepted with the acceptance probability $\min\{1, A_{MS1}\}$. Furthermore, if the state of the newly combined region is equally equal to the state of the adjacent region, we automatically merge these regions with the same CN state as a double merge. Two situations require a double merge. First, the $m$ and $m+1$ regions are merged into region $m^*$, and the state of the combined region is selected from $m$. If the state of the $m+2$ region is equal to that of $m^*$, we merge the $m^*$ and $m+2$ regions into the new region $m^{**}$. Then, the accepted probability is $\min\{1, A_{MS2}\}$. Second, the $m$ and $m+1$ regions are merged to region $m^*$, and the state of the combined region is selected from $m+1$. If the state of the $m−1$ region is equal to that of $m^*$, we merge the $m^*$ and $m−1$ regions into the new region $(m−1)^{**}$. Then, the accepted probability is $\min\{1, A_{MS3}\}$.

For the reverse strategy named split, one window boundary with a value of 0 (i.e., not a CN region breakpoint) is updated to a value of 1, and then, the CN region is split into two regions. The state of one newly formed region is randomly set to be the same as that of the original region, and the other region is restricted to be unequal to the states of the original and adjacent regions. Assume the selected window boundary belongs to the CN region $m$ and that the region $m$ is split to $m^*$ and $m^{**}$. Then, the accepted probability is $\min\{1, A_{RS}\}$.

Moreover, the reverse strategy of double merge, which is named trifid (split into three), essentially changes the values of two of the window boundaries with 0 values in one CN region (assuming the $m^{th}$ CN region) to 1 values, and then, three CN regions (indexed as $m^*$, $m^{**}$, and $m^{***}$) are constructed. The states of the leftmost ($m^*$) and rightmost ($m^{***}$) regions are assigned to be the same as that of the original region, and the state of the middle region ($m^{**}$) is randomly selected from the other states with equal probability. The accepted probability is $\min\{1, A_{TR}\}$.

Finally, the breakpoint of the CN region randomly shifts to the left or right window boundary with equal probability without changing the CN states for the boundary change strategy. The accepted probability of the left and right shift are $\min\{1, A_{RL}\}$ and $\min\{1, A_{RR}\}$, respectively. All of the above acceptance probabilities are derived in Supplementary Text 2 (Acceptance probabilities).

For setting the initial values of $C$ and $B$ in RJMCMC, a cubic smoothing spline model is fitted to the ordered read-depth signals (RDSs) across the windows. If the predicted RDSs in adjacent windows $i$ and $i+1$ are crossed to the threshold, then the window breakpoint $B_i$ is initially set to 1. The thresholds are approximately defined as the 5th and 95th percentiles of all predicted RDSs. According to the initial breakpoints, we randomly assign the initial state of each region, but the restriction of neighboring regions with different states should be satisfied. The probabilities of selecting four jumping strategies for updating $B$ are set as (1/3, 1/6, 1/6 and 1/3).

Additionally, to reduce the unbalanced effect that results from the extreme normal/abnormal state proportion, the whole genome is partitioned into several nonoverlapping sections to estimate the parameters. In our proposed procedure, we run RJMCMC for one genomic section at a time but set the initial values and hyperparameters based on whole genome to ensure that the evidence is sufficient. Advice regarding the section length is provided in the Results section.

Identification of copy number variations. The samples generated from the posterior distribution through RJMCMC are summarized to identify the CN states of windows via Bayesian testing statistics and Bayes factor (BF)\cite{36}. After burn-in ($t_{burn} = 5,000$ iterations, as the default setting), $K−1$ types of BF$s$ representing the strength for the abnormal states (CN = 1, 3, 4, …, $K$) against the normal state (CN = 2) in each window are calculated. The BF of window $i$ at iteration $t$ with abnormal CN state $k$ is defined as

$$P(C|D, B) \propto \prod_{m=1}^{M} \left[ \frac{1}{\sqrt{n_{cm}} \Gamma(n_{cm} + \frac{k}{2})} \beta_{cm}^{n_{cm} + \frac{k}{2}} \frac{1}{\Gamma(n_{cm})} \beta_{cm}^{-n_{cm}} \cdot \beta_{cm}^{n_{cm} + \frac{k}{2}} \right] \times \prod_{k=1}^{K} \left[ \frac{1}{\Gamma(1 + n_{rk})} \prod_{k'=1}^{K} \frac{1}{\Gamma(1 + n_{rk'})} \prod_{k'=1}^{K} \frac{1}{\Gamma(1 + n_{rk'})} \right]$$
The searching criteria for DGV were as follows: variant type each tool. The CNVs identified via each tool were compared with the summarized lists in the Database of Genomic Variants.

Results

Application to samples from the 1000 Genomes Project. For NA12156 and NA12878, after the preprocessing steps, approximately 220 megabasepairs (Mb) on chromosome 1 remained for the subsequent analysis. In CONY, approximately 440 sections with 0.5 Mb each were operated in parallel for RJMCMC sampling. The number of possible CN statuses was assigned as 5 (CN 1, 2, 3, 4, and 5) for the single-sample analysis and 3 (copy loss, normal, and copy gain) for the paired-samples analysis. The other parameter settings followed the default settings (see Supplementary Text 1). Some commonly used tools based on read depths (with hundreds of citations through March 2020) were compared. The competing algorithms (CNVnator, FREEC, and rdxplorer) used the default settings of 1, 3, 4, 5, 10, 25, 50, 100, 250, 500, and 1000 kb. The detection rate represents the recall for CNV regions, which is the percentage of CNV regions in the DGV that have any position identified as a CNV via the algorithm. The CNV region precision and base precision. The base recall is defined as the percentage of basepairs not listed as CNVs in the DGV that are also identified by the algorithm. The CNV detection rate represents the recall for CNV regions averaged over 100 case samples (for the single-sample analysis) or 1000 case-control pairs (for the paired-samples analysis).

In the 1000 Genomes Project analysis, the base accuracy is assessed by three numerical measurements, including the base recall (also called sensitivity), base false positive rate (FPR) and base precision. The base recall is defined as the percentage of basepairs classified as copy losses or gains. In addition, the CNV detection rates are calculated for each combination of 2 CNV types (copy loss/gain) for the paired-samples analysis. The other parameter settings followed the default settings (see Supplementary Text 1). Some commonly used tools based on read depths (with hundreds of citations through March 2020) were compared. The competing algorithms (CNVnator, FREEC, and rdxplorer) used the default settings of 1, 3, 4, 5, 10, 25, 50, 100, 250, 500, and 1000 kb. If the artificial CNV region is partially or fully identified, then the region is counted. Then, the detection rate is the percentage of detected artificial CNV regions averaged over 100 case samples (for the single-sample analysis) or 100 case-control pairs (for the paired-samples analysis).

For the simulation study, the base accuracy includes the overall base accuracy, base recall and base FPR. The overall base accuracy is summarized from the correctly identified basepairs. The base recall is defined as the percentage of CNV basepairs that are detected correctly. The base FPR is determined by the percentage of normal basepairs that is classified as copy losses or gains. In addition, the CNV detection rates are calculated for each combination of 2 CNV types (copy loss/gain) versus 10 CNV sizes (1, 2.5, 5, 10, 25, 50, 100, 250, 500 and 1000 kb). If the artificial CNV region is partially or fully identified, then the region is counted. Then, the detection rate is the percentage of detected artificial CNV regions averaged over 100 case samples (for the single-sample analysis) or 100 case-control pairs (for the paired-samples analysis).

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The CNVs identified via each tool were compared with the summarized lists in the Database of Genomic Variants. The searching criteria for DGV were as follows: variant type each tool. The CNVs identified via each tool were compared with the summarized lists in the Database of Genomic Variants. The searching criteria for DGV were as follows: variant type each tool.
**Table 1.** Performance of CNV detection in the experimental data analysis for NA12156 and NA12878. aThe number of CNV basepairs in the DGV that have any position identified as a CNV via the algorithm. bThe number of CNV regions in the DGV that identify at least one CNV basepair as a CNV.

| Sample(s)   | Algorithm | CNV bases | CNV regions | Detection rate |
|-------------|-----------|-----------|-------------|----------------|
| Single-sample analysis (NA12156) | CONY | 37,984 | 91.34% | 9.71% | 1.68% | 33 | 91.67% |
| | CNVnator | 343,308 | 84.30% | 13.07% | 1.15% | 25 | 69.44% |
| | FREEC | 86,204 | 21.17% | 13.19% | 0.34% | 1 | 5.56% |
| | rdxplorer | 284,865 | 69.95% | 2.28% | 5.27% | 11 | 30.56% |
| | DGV | 407,253 | 35.4X | 0.3% | 0.5% | 9 | 23.33% |
| Paired-samples analysis (NA12878) | CONY | 376,510 | 73.10% | 0.74% | 18.55% | 23 | 63.89% |
| | CNVnator | 163,695 | 31.78% | 15.44% | 0.47% | 29 | 80.56% |
| | FREEC | 378,282 | 34.61% | 6.63% | 1.18% | 3 | 8.33% |
| | rdxplorer | 515,073 | 91.43% | 0.51% | 1.65% | 25 | 83.33% |

**Table 2.** Performance comparisons in the simulation study.

| (%) | Algorithm | Overall accuracy | Recall | FPR | Precision |
|-----|-----------|------------------|--------|-----|-----------|
| Single-sample analysis | CONY | 99.21 | 99.44 | 0.70 | 97.85 |
| | CNVnator | 99.09 | 99.67 | 0.88 | 99.11 |
| | FREEC | 98.62 | 91.77 | 0.36 | 92.79 |
| | rdxplorer | 97.12 | 93.67 | 0.59 | 82.18 |

| | Algorithm | Overall accuracy | Recall | FPR | Precision |
|-----|-----------|------------------|--------|-----|-----------|
| Paired-samples analysis | CONY | 99.86 | 99.50 | 0.02 | 98.82 |
| | CNVnator | 94.65 | 71.10 | 0.01 | 47.35 |
| | FREEC | 99.72 | 98.49 | 0.06 | 98.90 |

Algorithm performance comparisons in a simulation study. The performance of the proposed procedure, CONY, was also compared with that of published methods for a single-sample analysis (CNVnator, FREEC, and rdxplorer) and paired-samples analysis (CNVSeq and FREEC) on simulation data. The competing algorithms utilized the default settings to identify the CNVs.

In the single-sample analysis, CONY performed satisfactorily in terms of overall base accuracy and base recall (Table 2). This comprehensive algorithm also had impressive CNV detection rates, especially for CNV sizes larger than 10 kb (Fig. 4a,b). The testing-based tool rdxplorer revealed great detection rates for all sizes of CNVs. However, the inaccurate breakpoints of the identified CNV regions yielded a low recall and high FPR. CNVnator was too rigorous to detect small CNVs (<10 kb), but its great performance in terms of the detection rates of the midsized and large CNVs contributed to its high overall base accuracy. Notably, CNVnator had high FPRs in detecting the absolute copy loss. FREEC had the worst performance in terms of the CNV detection rates for all sizes among all comparative methods. Overall, the methods had relatively high FPRs in deletion detection compared with duplication detection since copy loss was easier to identify than gain in sequencing platforms5.
In the paired-samples analysis, CONY was superior to the other methods in terms of CNV detection rates. While FREEC had slightly greater duplication recall than CONY, FREEC was significantly worse at finding small CNVs. CNVseq had a limited ability to detect CNVs.

In summary, CONY can detect both absolute and relative CNVs in single- and paired-samples analyses. CNVs with moderate to large sizes (>10 kb) can almost completely be detected by CONY. However, detecting small CNVs using a read-depth-based algorithm, including CONY, is challenging. The detection rates of small CNVs can be greatly improved by increasing the read coverage, which is demonstrated in the following results. Due to the poor power for detecting small CNVs for low-read-coverage data (e.g., 2.2X in our simulation), we suggest focusing on detecting CNVs with sizes >1,000 bp (as per the usual definition) to reduce potential false positives.

All simulations were run via the supercomputer Advanced Large-scale Parallel Supercluster (ALPS) at the National Center for High-performance Computing, National Applied Research Laboratories, Taiwan, with an AMD Opteron 6174 2.2 GHz × 4 CPU, a DDR3 ECC 128 GB of memory, and 512 nodes. In the RJMCMC procedure, one chromosome was divided into several nonoverlapping sections of equal size 0.5 Mb, and the operations were performed in parallel. The running time corresponded to the components of the CN in each analytic section. If only one CN state was included in the section, then the computing time was less than 1 minute. For a section with complex CN components, in our experience, the greatest length of time until RJMCMC became stable was less than 10 minutes. The running time for the other competing approaches with complex CN components are shown below: rdxplorer (~4 minutes), CNVnator (~15 minutes), FREEC (10 to 20 minutes), and CNVSeq (2 to 3 hours).

Analytic section length decision. To address the unbalanced structure of normal/variant regions in the genome, the whole genome can be partitioned into several nonoverlapping sections to estimate the parameters. The optimal section length for RJMCMC was derived via simulation. The samples generated for the algorithm comparisons in the above section were used. Six analytic lengths were adopted, including 60, 10, 5, 1, 0.5, and
0.1 Mb per section. Both the CNV detection rate and the base accuracy were used to select the proper section lengths.

Supplementary Fig. S2 presents the CNV detection rates using various CNV sizes and section lengths. As shown in the figure, the detection rates of CNVs of various sizes were enhanced by reducing the analytic section lengths. However, enhancing the detection rate appreciably for small CNVs (≤10 kb) was challenging, even after shrinking the section lengths. CNVs larger than 10 kb were considered to select an optimal section length. If the minimum requirement of the detection rate was set as 80%, then the section length should be shorter than 0.5 Mb. If a more severe detection rate was set, then a shorter section size was needed. In terms of CNV detection ability, the optimal section size was considered to be less than 0.5 Mb.

For the base accuracy, the results are shown in Supplementary Table S2. The recall was improved by shortening the section lengths. However, the FPRs dramatically increased when the sections were too small to provide sufficient evidence. In terms of the overall base accuracy, approximately 0.5–1 Mb (for single-sample analysis) and 0.1–0.5 Mb (for paired-samples analysis) were the proper section lengths for achieving peak accuracy. Based on the two performance measurements mentioned above, the recommendations for the section length were simplified to 0.5 Mb, which was also adopted in our experimental data analysis and simulation studies.

**Window read-depth estimation.** In this study, an alternative method was adopted for window read-depth estimation to enhance the completeness of the genetic information. Traditionally, the middle or start position of a read is located in a specific window, and the read is counted for the depth of this window. However, this strategy might underestimate the contribution of reads that span many windows. In our procedure, a summation approach was used. The read depths of each base were generated using the piling procedure in SAMTools, and then, the base depths in the specific window were summed as the window read depth. Supplementary Table S3 provides evidence that the summation method can improve both the CNV detection rate and the overall accuracy compared with the traditional representative-position method in single-sample analyses, especially for low coverage sequencing.

**Read coverage.** Because the NGS platform is still more expensive than other available technologies, researchers might process several samples in a single experimental run, which can result in low coverage. The depths based on sparse read coverage may lead to insufficient evidence for CNV identification. To evaluate the coverage effect, we followed the simulation settings mentioned above and generated 100 cases that were sequenced with a high coverage (22×). The CNV detection rates and base accuracies in the single-sample analysis are listed in Supplementary Table S3. Obviously, a great improvement was achieved in terms of CNV detection capability with high-coverage sequencing, especially for the detection of small variants. The impressive detection rates and outstanding recalls were attributed to the sufficient data information, but the false discoveries are expected to be accompanied by additional variations. Notably, no obvious differences were observed in the overall base accuracies between the low- (99.21%) and high-coverage (98.74%) data by CONY.

Two experimental samples (HG00419 and HG01595) from the 1000 Genomes Project, which were sequenced with both low (5.2 to 9.8X) and high (33.6 to 35.4X) coverages, were also analyzed to evaluate the coverage effect (Supplementary Table S1). High-coverage sequencing generally achieved better base accuracy and CNV detection rates in both single-sample and paired-samples analyses than low-coverage sequencing did for all tested algorithms. The base recall from CONY in the single-sample analysis is an exception, where high-coverage sequencing did not do better than low-coverage sequencing.

**Discussion**

Based on a comprehensive Bayesian hierarchical model and an efficient RJMCMC inference algorithm, the procedure proposed in this article was proven to be robust and precise for CNV detection. This functional tool can be applied for different purposes, including the detection of absolute and relative CNVs under single-sample and paired-samples designs. Samples from the 1000 Genomes Project were analyzed. CONY detected more CNVs and positions validated by the DGV database than the compared algorithms. In the simulation studies, the estimation methods performed well in terms of the overall base accuracy, recall and FPR for both single-sample and paired-samples analyses. Additionally, the CNV detection rates were effectively improved by selecting the proper analytic section length in the RJMCMC method and by adopting summation window read-depth estimation. The detection rates for small CNVs were still restricted even with suitable section lengths and depth estimation. In addition, we showed that the detection of small CNVs can be greatly improved by increasing the read coverage.

Although whole genome sequencing (WGS) is a comprehensive platform for exploring potential variants, target exome sequencing (TES) is an efficient choice because human exons constitute approximately 1% of the total genome but over 85% of genomic disease-causing regions. Exome sequencing provides effective information with high coverage on a limited budget. Read generation with WGS and TES follows distinct procedures due to the concentrations of DNA, the environments of hybridization and the methods of sequencing. Because of these experimental differences, the algorithms used to detect CNVs from WGS and TES are distinct, with alternative preprocessing, bias corrections and model assumptions.

WGS can detect more CNVs and precise breakpoints due to the complete genome scanning. WGS-based methods consider the entirety of the genome space, and the CNVs are estimated from the read depths across the genome with few significant bias corrections, such as for potential PCR duplicates and GC content. In contrast, the prediction of exact CNV breakpoints and small CNVs by segmentation algorithms in interrupted target exome sequences is challenging. In addition, exon-specific biases, such as exon sizes and batch and background effects, need to be corrected via multiple sample comparisons and/or additional adjustment steps. Therefore, the existing methods of WGS and TES seldom have commonalities. Modifying our approach for both WGS and TES under a common model framework will be a challenge for future research.
Data availability

The datasets used and analyzed in this study are available from 1000 Genomes Project (http://www.1000genomes.org). R code that implements the proposed procedure is available at https://github.com/weiyuchung/CONY, with direct links for downloading available at https://github.com/weiyuchung/CONY/archive/master.zip.

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Author contributions
G.-H.H. and Y.-C.W. conceptualized the research. Y.-C.W. developed the methods, built the CONY package, and performed data analysis. Y.-C.W. wrote the original draft, and G.-H.H. reviewed and edited the manuscript. G.-H.H. supervised the project. All authors read and approved the final manuscript.

Competing interests
The authors declare no competing interests.
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