| **Title**     | Proteome analysis of human metaphase chromosomes |
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| **Citation** | Journal of Biological Chemistry. 2005, 280(17), p. 16994-17004 |
| **Version Type** | VoR |
| **URL**      | https://hdl.handle.net/11094/79049 |
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| **Note**     | |
Proteome Analysis of Human Metaphase Chromosomes*

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DNA is packaged as chromatin in the interphase nucleus. During mitosis, chromatin fibers are highly condensed to form metaphase chromosomes, which ensure equal segregation of replicated chromosomal DNA into the daughter cells. Despite >1 century of research on metaphase chromosomes, information regarding the higher order structure of metaphase chromosomes is limited, and it is still not clear which proteins are involved in further folding of the chromatin fiber into metaphase chromosomes. To obtain a global view of the chromosomal proteins, we performed proteome analyses on three types of isolated human metaphase chromosomes. We first show the results from comparative proteome analyses of two types of isolated human metaphase chromosomes that have been frequently used in biochemical and morphological analyses. 209 proteins were quantitatively identified and classified into six groups on the basis of their known interphase localization. Furthermore, a list of 107 proteins was obtained from the proteome analyses of highly purified metaphase chromosomes, the majority of which are essential for chromosome structure and function. Based on the information obtained on these proteins and on their localizations during mitosis as assessed by immunostaining, we present a four-layer model of metaphase chromosomes. According to this model, the chromosomal proteins have been newly classified into each of four groups: chromosome coating proteins, chromosome peripheral proteins, chromosome structural proteins, and chromosome fibrous proteins. This analysis represents the first compositional view of human metaphase chromosomes and provides a protein framework for future research on this topic.

Chromosome formation during mitosis ensures the equal and appropriate segregation of genomic information into daughter cells (1). This surprisingly well organized process was first precisely described by Flemming in 1882 (2), and its structure has been investigated by numerous researchers. A metaphase chromosome consists of two DNA molecules and chromosomal proteins that are further divided into histones and non-histone proteins (1, 3). Histones and their structural roles have been well characterized, including their post-translational modifications (3–8). Several studies were also carried out to elucidate the vital role of non-histone proteins in chromosome structure and function. Laemmli and co-workers (9–11) and Earnshaw and co-workers (12) carried out pioneering work, and using electron microscopy in combination with extensive biochemical analyses, they suggested the existence of a proteinaceous chromosome scaffold to which the DNA is tethered in a loop formation based on their observation of histone-depleted chromosomes. Two proteins (topoisomerase IIα and SMC2) that constitute the scaffold were discovered through these studies (11, 12), although the contribution of topoisomerase IIα to the metaphase scaffold is still controversial (13–16). Several attempts have also been previously made to identify chromosomal proteins (17–20), and chromosome passenger proteins and proteins located at the chromosome periphery have been reported. In addition, some specially localized proteins in centromeric and telomeric regions have been reported (21–23). During the last decade, significant progress using mitotic extracts from frog eggs led to the identification of a condensin complex as an essential factor in chromosome condensation (24, 25). Despite the extensive investigation on the chromosomal proteins in the past, a fundamental question remains still unanswered. What is the global protein composition of the chromosome? This situation originates from the experimental difficulty in isolating intact chromosomes in large quantity for protein analyses given that chromosomes are formed only in a short period during the mitotic stage of the whole cell cycle in a few dividing cells.

To answer this question, the identity and localization of proteins constituting the chromosomes must be solved. Recent mechanical analyses of chromosomes using micromanipulators also raise questions regarding the scaffold model and would benefit from a list of the constituent proteins of metaphase chromosomes (26, 27). Recently, we reported several isolation methods for human metaphase chromosomes in large quantities after cell cycle synchronization and preliminarily compared them with each other in terms of their morphologies and protein compositions (28, 29). In this study, we used comparative proteomics and localization analyses to form a complete view of the chromosome from the protein perspective. We performed proteome analyses on three types of isolated human metaphase chromosomes. First, polyamine (PA)1 chromosomes,
isolated by the PA procedure, a standard preparation method for chromosomes both in biochemical and morphological studies (9, 28–30), were investigated in this study. Second, sucrose gradient (SG) chromosomes, purified by sucrose density gradient centrifugation (SDGC) of PA chromosomes, were prepared (29, 30). The proteins extracted from these preparations were separated by one-dimensional SDS-PAGE, followed by Coomassie Brilliant Blue staining. Quantitative information on proteins with molecular masses <100 kDa was obtained by isoelectric focusing (IEF) and radical-free and highly reducing (RFHR) (29, 31) two-dimensional gel electrophoresis. Electropherically separated proteins were identified by mass spectrometry-based proteomics. Finally, we prepared purified metaphase chromosomes, referred to as Percoll gradient (PG) chromosomes, which were purified by glycerol and Percoll density gradient centrifugation (9, 32). Subsequently, the localizations of several representative proteins on the chromosomes were assessed by indirect immunostaining. The identified proteins were newly classified into four groups based on their compositional characterization and localization.

**EXPERIMENTAL PROCEDURES**

**Reagents for Proteome Analysis—**Water was purified using a Milli-Q system (Millipore Corp., Bedford, MA). Digitonin was purchased from Sigma and EMD Biosciences. Water for in-gel digestion and mass spectrometry, acetonitrile, and acetic acid were purchased from Nacalai Tesque (Kyoto, Japan). Trifluoroacetic acid, Coomassie Brilliant Blue R-250, and lysyl endopeptidase were purchased from Wako (Osaka, Japan). Sequence-grade trypsin and endoproteinase Glu-C were purchased from Roche. Proteins used for molecular mass references and molecular mass determinations were purchased from Sigma and EMD Biosciences. Water was purified using a Milli-Q system. All other reagents were purchased from Wako or Sigma.

**Isolation of Human Metaphase Chromosomes—**PA chromosomes were isolated from synchronized human cell line (BALL-1) using a PA procedure, which was previously published (28) with minor modifications. The PA chromosomes were diluted in PA buffer (15 mM Tris-HCl (pH 7.2), 2 mM EDTA, 80 mM KCl, 20 mM NaCl, 0.5 mM EGTA, 0.2 mM spermine, 0.5 mM spermidine) and collected by centrifugation at 1750 × g for 10 min. This washing procedure was repeated three times. Although chromosomes maintain their condensed structure with 0.2 mM spermine, and 0.5 mM spermidine) and collected by centrifugation at 3000 × g for 30 min in a JS-24.38 rotor. A band containing chromosomes located at a level that was one-fifth the length of the tube measured from the bottom (Fig. S1) was diluted 3-fold in isolation buffer (5 mM Tris, 20 mM KCl, 30 mM EDTA, 0.25 mM spermidine, 1% thiodiglycol, 0.1 mM phenylmethylsulfonyl fluoride, and 0.1% Empigen) and subsequently centrifuged at 3000 × g for 15 min to collect PG chromosomes.

**Extraction of Proteins from Isolated Chromosomes—**For one-dimensional SDS-PAGE analysis, isolated chromosomes were directly suspended and dissolved in SDS-PAGE buffer (62.5 mM Tris, pH 6.8), 5% 2-mercaptoethanol, 2% SDS, and 0.05% bromophenol blue). For two-dimensional gel electrophoresis, proteins were extracted from isolated chromosomes using the acetic acid extraction method described previously (29, 34). Typically, a single extraction experiment using PA chromosomes isolated from 2.8 × 10^5 BALL-1 cells yielded 500 μg of proteins, whereas PG chromosomes isolated from 2.8 × 10^5 HeLa S3 cells contained 44 μg of proteins. The lower amount of proteins extracted from PG chromosomes originated mainly from the decrease in the chromosomes at each purification step. Protein aliquots were lyophilized and stored at −80 °C until further analysis.

**Electrophoretic Separation of Chromosomal Proteins—**For one-dimensional SDS-PAGE using 5–10 or 5–20% gel, Perfect NT gel (DRC Corp., Tokyo, Japan) was used. In addition to conventional IEF two-dimensional gel electrophoresis, RFHR two-dimensional gel electrophoresis was also used for the separation of chromosomal proteins covering a wide range of isoelectric points (29, 31). RFHR two-dimensional gel electrophoresis provides an excellent separation of highly basic proteins. The detailed procedures involved in IEF and RFHR two-dimensional gel electrophoresis analyses were described in our previous study (29).

**In-gel Digestion—**Pieces of the gel containing proteins of interest were excised for digestion with trypsin, lysyl endopeptidase Lys-C, or endoproteinase Glu-C using the following procedures. Gel pieces obtained from one- or two-dimensional gel electrophoresis were destained by incubation in 1 ml of 50 mM ammonium bicarbonate in 50% methanol. For reductive alkylation, gels dehydrated with 100% glycerol were excised for digestion with trypsin, lysyl endopeptidase Lys-C, or endoproteinase Glu-C from Roche. Proteins separated by one-dimensional SDS-PAGE were stained with Coomassie Brilliant Blue and then quantified using ImageQuant (Amersham Biosciences). In addition, the quantitative information on proteins with a molecular mass lower than ~100 kDa was mainly obtained from two-dimensional gel electrophoresis analyses. Proteins separated by two-dimensional gel electrophoresis were identified by ImageMaster 2-D Elite (Amersham Biosciences). The amount of each protein was determined as a ratio to core histone H4, which was well separated from the other core histones. The molar ratios of each protein in relation to histone H4 were calculated from the results obtained with the smallest possible starting amounts of protein to avoid artifactual underestimation resulting from stain saturation of the bands or spots in one-dimensional SDS-PAGE or two-dimensional gel electrophoresis, respectively. Thus, under these conditions, the amount of protein has a linear relationship with the intensity of the corresponding band or spot. The relative amounts of each protein were determined by averaging more than six two-dimensional gel electrophoresis results obtained under the same conditions. In the case that several proteins were identified from a single band, the band intensity was divided by the averaged molecular masses of the identified proteins, which provided the summation of the relative molar ratio of the identified proteins.
Proteome Analysis of Human Metaphase Chromosomes

gathered, followed by reduction of the volume to 5 μl using the SpeedVac. Desalting of the peptides was performed with ZipTip C₁₈ pipette tips (Millipore Corp.). After equilibration with 0.1% trifluoroacetic acid, peptides were adsorbed to the ZipTip C₁₈ pipette tips, followed by off-loading washing with 0.1% trifluoroacetic acid and elution with 5 μl of 50% acetonitrile containing 0.1% trifluoroacetic acid.

**Protein Identification by Mass Spectrometry**—For identification using the peptide mass fingerprinting (PMF) method, 1-μl aliquots were deposited on an AnchorChip 384-well target plate (Bruker Daltonik GmbH, Bremen, Germany) followed by deposition of 1 μl of matrix (10 mg/ml-cyano-4-hydroxycinnamic acid in 50% acetonitrile and 0.1% trifluoroacetic acid). Subsequently, the drops were dried up in the atmosphere. The identification of each protein after the enzymatic digestions was carried out by PMF analysis using a matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) mass spectrometer (Autoflex or Ultraflex, Bruker Daltonik GmbH) and by tandem mass spectrometry (MS/MS) analysis using 1 MALDI-TOF/TOF mass spectrometer (Ultraflex) (35). In addition, an Esquire electrospray ionization MS/MS system (Bruker Daltonik GmbH) equipped with a liquid chromatography (Dionex Corp., Sunnyvale, CA) was employed for protein identification. Mass data extracted by Flex analysis (Bruker Daltonik GmbH) for MALDI-TOF/TOF or by Hystar Version 2.3 (Bruker Daltonik GmbH) for electrospray ionization data were analyzed using Bioteqs Version 2.2 (Bruker Daltonik GmbH) employing the MASCOT search engine. Most proteins were identified by matching the PMF results with the NCBIin Data

**Immunofluorescence Microscopy**—Monoclonal antibodies against β-actin and histone H1 were produced for this study. The production of monoclonal antibodies against these chromosomal proteins was carried out by immunization of mice with the extracted chromosomal proteins. After screening by indirect immunofluorescence microscopy, several monoclonal antibodies were obtained. The immunoprecipitated antigen protein was identified by PMF.² Polyclonal antibody against human chromosome-associated protein (hCAP) G was produced as described previously (36). Antibodies against BIP and hSNF2H (H-300) were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Monoclonal antibody against Ki-67 (MIB-1) was purchased from Dako Cytomation (Glostrup, Denmark). Antibody against calreticulin was purchased from Alexis Biochemicals (Lausen, Switzerland).

Indirect fluorescent antibody staining of HeLa cells was performed according to a previously published method (37). HeLa S3 cells were grown on coverslips coated with poly-l-lysine (Sigma). The cells were according to a previously published method (37). HeLa S3 cells were purchased from Alexis Biochemicals (Lausen, Switzerland). Antibody against calreticulin was produced as described

**RESULTS**

**Morphologies and Protein Compositions of Isolated Human Metaphase Chromosomes**—PA chromosomes are free of nuclei, as previously confirmed by flow cytometry (28). Optical microscopy demonstrated that PA (29) and SG (Fig. 1A) chromosomes retained the native morphology of metaphase chromosomes. One-dimensional SDS-PAGE of PA chromosomes (Fig. 2, A and B) revealed numerous protein bands spanning a broad molecular mass range in addition to reasonably intense bands of the most abundant proteins, core and linker histones. The weight percentage of histones in all of the chromosomal proteins was estimated at 60%, 48 and 12% of which correspond to core and linker histones, respectively. One-dimensional SDS-PAGE indicated that the DNA-dependent protein kinase catalytic subunit (465 kDa) had the highest molecular mass. Two intense bands corresponding to myosin II and a 190-kDa leucine-rich protein were confirmed by one-dimensional SDS-PAGE in the higher molecular mass region (Fig. 2B, bands 4 and 9). The IEF two-dimensional gel electrophoresis patterns of PA and SG chromosomal proteins at pl 4–7 and 6–11 are shown in Fig. 3. The RFHR two-dimensional gel electrophoresis results for PA and SG chromosomal proteins are provided in Fig. 4 (A and B, respectively). PA chromosomes had a larger number of proteins compared with SG chromosomes, especially in the weak basic regions, which are indicated in Fig. 3 (C and D, dotted ovals). Over 200 and 150 spots were reproducibly detected for PA and SG chromosomes, respectively. A chromatin-remodeling leucine-rich protein (spot A1; molecular mass of 145 kDa) had the highest molecular mass and was well separated at pl 4–7 (Fig. 3, A and C).

**Identification of Proteins in PA and SG Chromosomes**—Most

![Fig. 1. Microscopic observations of isolated human metaphase chromosomes.](image347x645 to 533x738)

A and B, Giemsa staining of SG chromosomes from BALL-1 cells and PG chromosomes from HeLa S3 cells, respectively.

mm KCl at 1.5–2.0 × 10⁵ cells/ml for 15 min. The cells were spread onto coverslips using a CytoSpin (Thermo Electron Corp.) and then fixed with 2% paraformaldehyde in PBS (pH 7.4) for 15 min at room temperature. After washing three times with PBS, the metaphase chromosome spreads were permeabilized in 0.2% Triton X-100 in PBS for 5 min at 4 °C. After washing, the spreads were incubated with 3% BSA in PBS for 30 min at room temperature. Subsequently, the spreads were incubated with primary antibodies and 3% BSA in PBS for 1 h at room temperature. After washing three times with PBS for 5 min, the spreads were incubated with secondary antibodies and 3% BSA in PBS for 1 h at room temperature. After being washed three times with PBS for 5 min, the spreads were mounted with 1 μg/ml 4′,6-diamidino-2-

phenylindole in Vectashield.

Isolated metaphase chromosomes were also immunostained according to a previously published method (37). Briefly, isolated chromosomes were diluted in XBE2 buffer and fixed by adding 0.1 volume of 8% paraformaldehyde at room temperature for 15 min. The fixed chromosomes were spun onto poly-l-lysine-coated coverslips through XBE2 buffer containing 30% sucrose and stained as described above for indirect fluorescent antibody staining of HeLa cells.

2 T. Higashi, S. Miyakawa, S. Uchiyama, S. Matsunaga, H. Takata, S. Fujimoto, M. Noda, A. Terauchi, T. Shimizu, M. Oda, T. Azuma, and K. Fukui, manuscript in preparation.
of the intense bands or spots observed after electrophoresis were unambiguously identified. In the PA and SG chromosomes, 209 proteins were identified, only 51 of which overlapped in one-dimensional SDS-PAGE and the two types of two-dimensional gel electrophoresis. Thus, 158 proteins were identified among the PA and SG chromosomes. Generally, their amounts estimated from one-dimensional SDS-PAGE were similar to those obtained by the two-dimensional gel electrophoresis analyses. The names of the proteins, their numbers corresponding to those in Figs. 2–4, database accession numbers, molecular masses, pI values, MASCOT scores, sequence coverage, Nuclear Protein Database accession numbers (39), and related information where available have been summarized (Tables S1–S3). The molar amount of each protein is provided per 100 histone H4 molecules. The differences in the molar amount of each protein between PA and SG chromosomes are indicated as molar ratios. One-dimensional SDS-PAGE provided quantitative information on proteins with molecular masses >100 kDa, whereas IEF two-dimensional gel electrophoresis provided information on acidic and basic proteins with molecular masses <100 kDa. RFHR two-dimensional gel electrophoresis provided complementary information on basic proteins. Proteins identified in PA and SG chromosomes were classified into six groups based on their known interphase localization. The six groups were nuclear, mitochondrial, ribosomal, cytoplasmic, cytoskeletal, and unknown.

**Protein Constituents of PA and SG Chromosomes—**Fig. 5A shows the percentages of classified proteins in PA chromosomes by number and molar amount. Mitochondrial proteins were the most abundant (37.7%, 60:159) with regard to number. Nuclear proteins represented 29.6% (47:159), and ribosomal proteins amounted to 13.2% (21:159) of the total proteins. Cytoplasmic proteins composed 12.0% (19:159). The molar amounts of the proteins estimated from the division of band or spot intensities by their molecular masses present an alternative view of the protein composition (Fig. 5A). Nuclear proteins represented 86.9% of the total molar amount of the PA chromosomal proteins. A conspicuous difference was due to a large amount of nuclear proteins and a small amount but substantial numbers of ribosomal, mitochondrial, and cytoplasmic proteins.
PA chromosomal proteins contained 34.3 histone H1 molecules/100 histone H4 molecules; this value is close to that reported for the interphase nucleus (40–42). The amount of high mobility group (HMG) N2 (Fig. 4A, spot R85) was 29.18 molecules/100 histone H4 molecules; and hence, it was the second most abundant protein following histones. HMGA1 (HMG AT hook protein-1) was present in a large amount (Fig. 2A, band 37; Fig. 3B, spot B30; and Fig. 4A, spots R81 and R83). Ubiquitinated histone H2A was also present in a large amount (Fig. 2A, band 34; Fig. 3B, spot B25; and Fig. 4A, spot R53).

Quantitative analyses based on the comparative proteome analysis between PA and SG chromosomes provided an important perspective for the identified proteins; they could be classified into two discrete types of protein groups with low and high affinities for isolated chromosomes (Fig. 5B). Proteins with low affinities were eliminated after SDGC (spots within the dotted ovals in Figs. 3C and 4B). For example, the mitochondrial 60-kDa heat shock protein was largely dissociated in SG chromosomes (Fig. 3A and C, spot A8). The amounts of most of the mitochondrial and cytoplasmic proteins became significantly low in SG chromosomes. The 21 proteins specific to PA chromosomes were mitochondrial proteins (Table S2). On the other hand, intense spots of histones, HMGs, and β-actin (Figs. 3 and 4) were similar in both PA and SG chromosomes, indicating their structural roles or high affinities for isolated metaphase chromosomes. Among nuclear proteins, the amounts of several proteins, including nucleolin, decreased, but a considerable amount still remained after SDGC.

Identification of Proteins in PG Chromosomes—For further identification and characterization of chromosomal proteins essential for the chromosome structure, proteome analyses of purified metaphase chromosomes (PG chromosomes) were performed by one-dimensional SDS-PAGE (Fig. 6) and RFHR two-dimensional gel electrophoresis (Fig. 7). Consequently, 107 major proteins involved in the detected bands or spots were identified (Supplemental Table S3). A sample of proteins identified in PA, SG, or PG chromosomes is given in Table I. Although 10 mitochondrial or cytoplasmic proteins were still identified from PG chromosomes, most of the mitochondrial and cytoplasmic proteins (e.g. typical mitochondrial or endoplasmic reticulum marker proteins, glutamate dehydrogenase or calreticulin, respectively) were absent in PG chromosomes. On the other hand, known chromosomal proteins such as topoisomerase IIα, condensin subunits, and inner centromere protein were frequently identified in PG chromosomes.

Localization of Representative Identified Proteins—The localizations on the chromosomes of representative proteins identified in this study were visualized by indirect immunostaining under three different cytological conditions in chromosome preparations. The three cytological conditions of isolated chromosomes in suspension (Fig. 8), chromosomes in cells (Fig. S2), and ordinary spread chromosome specimens (Fig. S3) would reflect isolated chromosomes, in vivo chromosomes, and the cytological control, respectively. Calreticulin was detected around isolated chromosomes and the cytological control, but not detected on in vivo chromosomes. This group included BiP. The other proteins except for those of this type were all localized on the chromosomes prepared under the three different conditions.
conditions. They were divided into two obviously different types. Ki-67 was visualized at the peripheral regions of two chromatids of all three types of chromosomes. Heterogeneous nuclear ribonucleoprotein (hnRNP) A2/B1 showed similar distribution patterns. Histone H1 and human SNF2H demonstrated rather uniform localization on all chromosomes. hCAP-G depicted axial localization at the midribs of both chromatids. β-Actin was distributed on chromosomes unevenly, with speckled intense signals at the periphery. Based on known information, mitochondrial and cytoplasmic proteins were clas-

Fig. 4. RFHR two-dimensional gel electrophoresis patterns of proteins from PA (A) and SG (B) chromosomes. The dotted oval in B represents the area in which the signal intensity of the spots markedly decreased after purification by SDGC.

Fig. 5. Profiles of the constituent proteins of PA and SG chromosomes. A, the identified proteins were classified based on their known localization during interphase. Percentages are presented based on either the number of the identified proteins (left) or the molar amounts of the identified proteins (right). B, differences in the amounts between PA and SG chromosomes are shown for the representative proteins from the four groups. The relative amounts of proteins in SG chromosomes are indicated as the ratios to those in PA chromosomes.
sified as one type of protein that was not localized on in vivo chromosomes with low affinity for isolated chromosomes. Furthermore, fibrous proteins such as \( \beta \)-actin, vimentin, and myosin consisted of one discrete type of protein.

**DISCUSSION**

To better define the protein constituent of metaphase chromosomes, the 157 proteins identified in PA and SG chromosomes and the 107 proteins in PG chromosomes were individually compared based on the following three criteria: protein behavior during SDGC and Percoll density gradient centrifugation, localization of each protein on metaphase chromosomes, and information on functional and biochemical properties. The protein behavior enabled us to classify the identified proteins into two types as indicated above. Based on the results, all of the identified chromosomal proteins were newly classified into the following four groups: 1) chromosome coating proteins (CCPs), 2) chromosome peripheral proteins (CPPs), 3) chromosome structural proteins (CSPs), and 4) chromosome fibrous proteins (CFPs) (Fig. 9A).

CCPs—The amounts of CCPs decreased markedly in SG chromosomes and were low in PG chromosomes. CCPs are mainly mitochondrial and cytoplasmic proteins found in small amounts (6.4%) but in a significant number (49.7%, 79:159) in PA chromosomes. Mitochondrial proteins with basic pI appear to be nonspecifically and weakly bound to chromosome surfaces during the chromosome isolation processes. The identified cytoplasmic proteins, including endoplasmic reticulum proteins, are mainly acidic. The 70-kDa heat shock protein is a typical acidic and weak binding protein that was largely removed by SDGC (Fig. 3, A and C, spot A4). The acidic proteins in PA chromosomes are likely to interact with the basic histones in the chromosome suspension. It appears that these proteins are not part of the chromosomes in living cells, where the subcellular organelles are maintained in their intact forms. The detergent used in the chromosome isolation causes the release of organelle proteins, thereby leading to their nonspecific binding to the surface of isolated chromosomes.

**Fig. 6. Profiles of the constituent proteins of PG chromosomes.** The proteins identified are indicated. The same number of proteins was also used in Table S3. INCENP, inner centromere protein; CENP-C, centromere protein C; PARP, poly(ADP-ribose) polymerase.

**Fig. 7. RFHR two-dimensional gel electrophoresis patterns of proteins from PG chromosomes.**
CPPs—Nuclear proteins have thus been classified into two new groups, viz. CPPs and CSPs. Proteins classified as CPPs are mainly nucleolar and nuclear envelope-related proteins. They were partly removed by SDGC. However, most of the CPPs that were identified in PA and SG chromosomes in significant amounts were also identified in PG chromosomes, thus indicating that they are the essential proteins for metaphase chromosomes. Proteins in this group were the second most significant amounts were also identified in PG chromosomes, thus indicating that they are the essential proteins for metaphase chromosomes. Proteins in this group were the second frequencies identified proteins (28%) in PG chromosomes. From the beginning of chromosome research, the existence of CPPs was visually detected, and they were referred to as the “chromosome matrix” or “chromosome sheath” (49). In mitotic pro-metaphase, the nucleolus disappears after the breakdown of the nuclear envelope (1). The existence of fibrillarin at the chromosome periphery has already been reported (1, 17). Other proteins such as hnRNP A2/B1 are components of the telomere complex (47) and Ku70 is involved in telomere maintenance (48); their localization in metaphase is not only in the telomeric region, but also in the entire peripheral region of metaphase chromosomes (49). This fact presents an interesting dynamic feature of telomere proteins in action. Presumably, they are pooled in the peripheral region as CPPs and function in the telomeric region of metaphase chromosomes.

Some ribosomal proteins were still found in SG and PG chromosomes, indicating the existence of certain ribosomal proteins that are different from ordinary ribosomal proteins classified as CPPs. This view is supported by reports in which ribosomal protein S1 was localized to the chromosome peripheral region (17) and ribosomal protein S6 was a component of ribosomal complexes (50).

CSPs—Because CSPs form the body of chromosomes and play a role in the condensation/decondensation of chromosomes (1), they should be common to PA, SG, and PG chromosomes. More than half of the number (57.5%, 27:47) of the nuclear proteins identified in PA chromosomes belonged to this group, and CSPs amounted to >80% of the PA chromosomal proteins (Fig. 9B). CSPs were most frequently (40.2%) identified in PG chromosomes (Tables S1 and S2). The proteins of the nuclear envelope in metaphase chromosomes among the three types of chromosomes analyzed.

### Table I

A sample of the proteins identified in PA, SG, and PG chromosomes

| No. | Protein                                      | pI  | Mass     | New classification | Localization at interphase |
|-----|----------------------------------------------|-----|----------|---------------------|---------------------------|
| 9, A1 | 130-kDa leucine-rich protein                 | 5.4 | 145,109  | CPPs                | Cy, Cs, N                 |
| 16, A4, R2, P19, r2 | 70-kDa heat shock protein 9B (mortalin-2)     | 6.0 | 73,882  | CPPs                | Cy                        |
| 20, A8, R6 | Chaperonin-60, Hsp60               | 5.7 | 61,016  | CPPs                | M                         |
| 2 | DNA-dependent protein kinase catalytic subunit | 6.9 | 465,266  | CPPs                | N                         |
| R4, r6 | Nucleolin                                 | 4.59 | 76,298  | CPPs                | N                         |
| A16, R19, r10 | hnrNP H                              | 5.9 | 49,198  | CPPs                | N                         |
| 23, B12, R25, P30, r16 | RNA-binding motif protein, X chromosome (hnRNP G) | 10.1 | 42,306  | CPPs                | N                         |
| 27, R32, P36, r23 | Fibrillarin                           | 10.2 | 33,407  | CPPs                | N                         |
| R47, r31 | Ribosomal protein S6                  | 10.9 | 28,614  | CPPs                | R                         |
| R17, r13 | Nucleophosmin/B23.2                  | 4.56 | 28,383  | CPPs                | N                         |
| 19, A8, r6 | Chaperonin-60, Hsp60               | 5.7 | 145,109  | CPPs                | Cs, N                     |
| 17, P17 | Lamin A/C                              | 8.6 | 70,618  | CPPs                | N                         |
| A16, R19, r10 | hnrNP H                              | 5.9 | 49,198  | CPPs                | N                         |
| 27, R32, P36, r23 | Fibrillarin                           | 10.2 | 33,407  | CPPs                | N                         |
| R47, r31 | Ribosomal protein S6                  | 10.9 | 28,614  | CPPs                | R                         |
| R17, r13 | Nucleophosmin/B23.2                  | 4.56 | 28,383  | CPPs                | N                         |
| 2 | DNA-dependent protein kinase catalytic subunit | 6.9 | 465,266  | CPPs                | N                         |
| R4, r6 | Nucleolin                                 | 4.59 | 76,298  | CPPs                | N                         |
| A16, R19, r10 | hnrNP H                              | 5.9 | 49,198  | CPPs                | N                         |
| 27, R32, P36, r23 | Fibrillarin                           | 10.2 | 33,407  | CPPs                | N                         |
| R47, r31 | Ribosomal protein S6                  | 10.9 | 28,614  | CPPs                | R                         |
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| R17, r13 | Nucleophosmin/B23.2                  | 4.56 | 28,383  | CPPs                | N                         |
| 2 | DNA-dependent protein kinase catalytic subunit | 6.9 | 465,266  | CPPs                | N                         |
| R4, r6 | Nucleolin                                 | 4.59 | 76,298  | CPPs                | N                         |
| A16, R19, r10 | hnrNP H                              | 5.9 | 49,198  | CPPs                | N                         |
| 27, R32, P36, r23 | Fibrillarin                           | 10.2 | 33,407  | CPPs                | N                         |
| R47, r31 | Ribosomal protein S6                  | 10.9 | 28,614  | CPPs                | R                         |
| R17, r13 | Nucleophosmin/B23.2                  | 4.56 | 28,383  | CPPs                | N                         |

* Numbers correspond to bands in one-dimensional SDS-PAGE and spots IEF and spots in (A, pI 4–7); and (B, pI 6–11) and RFHR (R and r) two-dimensional electrophoresis analyses.

* N, nucleus; M, mitochondrial; Cy, cytoplasmic; Cs, cytoskeleton; r, ribosome.
histone, ubiquitinated H2A (52), were also identified in all three types of isolated chromosomes. Significant amounts of macroH2A (Fig. 2A, band 25; and Fig. 6, band P33) were expected because this variant is known to localize on chromosomes during mitosis, overlapping with histone H3 methylated at Lys4 (53). HMGN2 and HMGA1 were the second and third most abundant proteins, respectively. These results suggest an important structural role of these particular HMG proteins, which is further supported by their high affinity for the chromatin fiber compared with other HMGs such as HMGB (54, 55). Poly(ADP-ribose) polymerase, hSNF2H, and the DEK oncogene protein were identified (Fig. 2, A and B). Inner centromere protein and Aurora B identified in PG chromosomes are known as chromosome passenger proteins and are localized to the centromeric region in metaphase (56). A new passenger protein recently reported, Borealin (57), also named Dasra B (58), was also identified as CDC8 in this study. Localization of poly(ADP-ribose) polymerase on the centromere, neocentromere, and chromosome arm has already been reported (59).

One of the features emerging from analyses using the three types of chromosomes was that not all components of the centromeric/kinetochoric and telomeric regions of chromosome were detected, although several proteins that concentrate in the centromeric region during mitosis, such as chromosome passenger proteins and heterochromatin protein-1 (60), were identified. Centromere protein C was the only protein among centromere proteins identified sporadically in this study. These facts reasonably suggest that centromere/kinetochore and telomere proteins are present only in small amounts compared with the overall chromosomal proteins. In fact, the quantitative purification of centromere protein A and telomeric repeat-binding factor-1 from human cells revealed markedly small molar amounts of ~1:70,000 and 1:100,000 histone H4, respectively (61, 62).

All eight subunits of the condensin I and II complexes (24, 25) were identified in PG chromosomes, although previous analyses of the human metaphase chromosome scaffold did not detect two of them, hCAP-D3 and hCAP-H2 (63, 64). Using a human mitotic cell extract, Takemoto et al. (65) reported that one condensin complex is present per 13 kilobase pairs of DNA. Assuming that one nucleosome is present in every 200 bp of DNA, this would imply that the molecular number of the condensin complex/100 histone H4 molecules is 0.77. Considering the molecular mass of hCAP-C (138 kDa) and that of histone H4 (11 kDa), the band intensity of hCAP-C should be approximately one-tenth that of histone H4. However, the band of hCAP-C in the one-dimensional SDS-PAGE pattern was considerably weaker than that expected from this estimation (Fig. 6, band P4). Based on the band intensity, the molar ratio of hCAP-C to 100 molecules of histone H4 is 0.32 at the most, which means they bind to chromatin fiber every 33 kilobase pairs on average. This discrepancy may be explained by the dissociation of the condensin subunits from metaphase chromosomes during the isolation procedures, despite the fact that
the isolated chromosomes retained the axial distribution of the condensin subunit (hCAP-G) in chromosomes (Fig. 8 and Figs. S2 and S3) and the native morphology when observed by optical microscopy. It has been reported that condensin subunits dissociate from metaphase chromosomes in the absence of divalent cations (9) or ATPase inhibitors (37).

**CFFs**—CFFs were classified primarily as an independent group because they are fibrous in nature and because their amounts were not changed in both PA and SG chromosomes, e.g. the spot intensity of β-actin did not change significantly (Fig. 3, A and C, spot A19). Mitotic kinesin-like protein, which interacts with the mitotic spindle, has also been classified into this group (66). Among 18 different CFFs, β-actin, vimentin, and tubulin were even identified in PG chromosomes (Fig. 6), which differs from a previous report (32). The contribution of CFFs to chromosome structure is still ambiguous (67). Myosin II was not identified in this study, although its localization in salamander nuclei (71). Furthermore, our observations obtained by indirect immunofluorescence microscopy suggested that metaphase chromosomes would play a definite structural role. Our observations obtained by indirect immunofluorescence microscopy indicated for the first time the localization of β-actin to metaphase chromosomes (Fig. 8). Mandeville and Rieder (73) have reported that metaphase chromosomes are surrounded by a “cage” of keratin filaments that restrict the dispersion of chromosomes during the breakdown of the nuclear envelope or microtubule penetration between and through the chromosomes. Thus, the CFFs that maintain the spatial organization of chromosomes within a mitotic cell are also likely to be bound to chromosomes in vivo.

**Candidates for Novel Chromosomal Proteins**—Fifteen proteins that were not classified into any of the four groups were identified in PA, SG, and/or PG chromosomes. To our knowledge, chromosomal localization of these proteins has not been reported before; thus, they could be novel chromosomal proteins.

**Conclusion**—We have provided a qualitative and quantitative identification of all of the major chromosomal proteins of isolated human metaphase chromosomes. Comparative proteome analyses of the chromosomes obtained by the three types of isolation procedures enabled us to differentiate chromosomal proteins as intrinsic chromosomal proteins or possible attachment proteins during the isolation procedures. The proteins identified here could be classified into four major different groups based on a combination of the results from the comparative proteome and localization analyses. A four-layer model of the metaphase chromosome has been developed in which most of the chromosomal proteins are allocated to one of the four layers; coating, peripheral, structural, and fibrous. Therefore, this study provides an important reference for future studies on chromosome structure and function.

**Acknowledgments**—We are grateful to Juan Ausio (University of Victoria) and Jordanka Zlateanova (University of Wyoming) for helpful discussions and critically reading the manuscript. We thank Akio Kobayashi and Eiichiro Fukusaki (Osaka University) for access to the liquid chromatography-MS/MS instrument and Akira Wada and Hideji Yoshida (Osaka Medical University) for technical help in RP-HPLC two-dimensional gel electrophoresis experiments. We thank Keiji Kimura (RIKEN Institute) and Fumio Hanaoka (Osaka University) for anti-hCAP antibodies. We also thank Sumire Inaga (Tottori University) for using the SEM picture of metaphase chromosomes with slight modification.
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