Epitope mapping of monoclonal antibodies specific for the 190-kDa multidrug resistance protein (MRP)

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Summary Inherent or acquired resistance to multiple natural product drugs in human tumour cells is often associated with increased expression of multidrug resistance protein (MRP), a 190-kDa integral membrane protein that belongs to the ATP-binding cassette (ABC) superfamily of transport proteins. Both clinical and experimental investigations of MRP have been facilitated by several monoclonal antibodies (MAbs) generated against intracellular epitopes of the molecule. Recently, however, several new ABC transporters that are quite closely related to MRP have been identified, raising concerns about the specificity of the MRP-reactive MAbs. In the present study, we have mapped the epitopes of MAbs MRP1 and MRPm6 to the decapeptides 239GSDLWSLNKE247 (located in the intracellular loop between the first and second membrane-spanning domains of MRP) and 1511PSDLLQRGRL520 (located near the carboxy terminus of MRP) respectively. Alignment of the MRP1 and MRPm6 epitope sequences with the comparable regions in mammalian ABC proteins most closely related to MRP indicates that, with the exception of murine mrp, the sequences are poorly conserved. We conclude that MAbs MRPm6 and MRP1, together with MAb QCRL-1, which has previously been mapped to the heptapeptide 8113SYSGD154, remain highly specific probes for detection of different regions of the MRP molecule.

Keywords: multidrug resistance; multidrug resistance protein; monoclonal antibody; epitope mapping

Successful treatment of many human tumours is often limited by the development of drug resistance. Experimentally, resistance to natural product drugs may be conferred by overexpression of one or other of the integral membrane proteins, MRP or P-glycoprotein, both of which are members of the ATP-binding cassette (ABC) superfamily of transport proteins (Deeley and Cole, 1997). ABC transporters typically consist of a hydrophobic polytopic membrane-spanning domain (MSD) followed by a cytosolic nucleotide-binding domain (NBD) that contains three signature motifs (Higgins, 1992). Many eukaryotic ABC proteins, including the 190-kDa MRP and 170-kDa P-glycoprotein, contain two MSDs and two NBDs organized in a tandemly duplicated fashion. However, unlike P-glycoprotein, MRP and several other closely related proteins contain an additional NH2-proximal MSD (Cole et al. 1992; Bakos et al. 1996; Loe et al. 1996a; Deeley and Cole, 1997; Stride et al. 1996). We have recently shown by site-directed mutagenesis of N-glycosylation sites that, in MRP, this MSD contains an odd number of transmembrane helices and, in contrast to P-glycoprotein, the amino-terminus of MRP is extracytosolic (Hipfner et al. 1997).

Whereas both MRP and P-glycoprotein confer resistance to chemotherapeutic agents such as doxorubicin and vincristine by reducing cellular accumulation of these drugs, MRP has also been demonstrated to be a primary active transporter of several structurally diverse conjugated organic anions. Known high-affinity substrates include the cysteinyl leukotriene (LT) C4, 17β-oestradiol 17-(β-d-glucuronide) and glutathione conjugates of activated aflatoxin B1 (Leier et al. 1994; Muller et al. 1994; Loe et al. 1996b, c, 1997). However, conjugation is not known to play an important role in the metabolism of most of the drugs included in the MRP resistance phenotype, and it has not been possible to demonstrate active transport of unconjugated substrates in vitro (Muller et al. 1994; Jedlitschky et al. 1996; Loe et al. 1996b), although we and others have demonstrated that MRP-mediated ATP-dependent transport of vincristine and aflatoxin B1 can occur in the presence of reduced glutathione (Loe et al. 1996b; Barnouin et al. 1997).

Numerous studies generally support the notion that P-glycoprotein plays a role in the drug resistance observed in several human malignancies (Filipits et al. 1996b). Although studied to a lesser extent, MRP also appears to be clinically important in a number of haematological and solid tumours, including neuroblastoma, certain instances of drug resistant retinoblastoma, some subtypes of non-small-cell lung cancer and breast cancer (Ota et al. 1995; Filipits et al. 1996a; Giaccone et al. 1996; Norris et al. 1996; Chan et al. 1997; Nooter et al. 1997).

To facilitate both experimental and clinical studies of MRP, several MRP-reactive monoclonal antibodies (MAbs) have been generated (Flens et al. 1994, 1996; Hipfner et al. 1994). However, the epitope of only one of these, MAb QCRL-1, has been mapped to single amino acid resolution (Hipfner et al. 1996). MAb QCRL-1 was raised against membranes from H69AR drug-resistant lung cancer cells, which express high levels of MRP. We have previously determined that the critical core of its epitope is the heptapeptide '6113SYSGD154' located in the cytosolic region linking the first NBD to the third MSD of the MRP molecule. Two other MRP-reactive MAbs, namely, rat MAb MPR1 and murine MAb MRPm6, were raised against fusion proteins, one containing...
a 167 amino acid peptide from the amino-proximal region of human MRP and the other containing a discontinuous 172 amino acid peptide from the carboxy-proximal region of the protein (Flens et al. 1994). Like MAb QCRL-1, these Mabs recognize linear intracellular epitopes of MRP and have been very useful for protein characterization and subcellular localization, as well as for immunohistochemical detection of MRP in normal and malignant cells and tissues (Filipits et al. 1996a; Flens et al. 1996; Nooter et al. 1995). In the present study, we have determined that the epitope sequences of Mabs MRP1 and MRP6 are localized to amino acids 238–247, located in the intracellular loop between the first and second MSDs of MRP, and amino acids 1511–1520, located carboxy-proximal to the second NBD of MRP very near to the carboxy terminus of the protein respectively.

**MATERIALS AND METHODS**

**Antibodies**

Rat MAb MRP1 (IgG₁) and mouse MAb MRP6 (IgG₁) were raised against maltose-binding protein fusion proteins containing human MRP amino acids 194–360 and human MRP amino acids 1294–1430 plus 1497–1531 respectively (Flens et al. 1994). Mouse MAb QCRL-1 (Cenocor Diagnostics, Malvern, PA, USA) was raised against cell membranes from MRP-overexpressing H69AR cells (Mirsik et al. 1987) and has been shown to bind to the heptapeptide —SSYSGDE— in the connector region of human MRP (Hijfner et al. 1994, 1996). MRP-1 and MRP-2 are rabbit polyclonal antisera raised against two 15 amino acid peptides containing the 'active transport' family signature sequences from the first and second NBDs of MRP (amino acids 765–779 and 1427–1441 respectively) (Hijfner et al. 1996, 1997).

**Generation of constructs, production of recombinant baculovirus and viral infection**

Two constructs encoding MRP molecules with NH₂-terminal deletions were prepared and transferred into the recombinant donor plasmid pFASTBAC1 (Life Technologies, Burlington, Ontario, Canada) as described elsewhere (Gao et al. 1998). These constructs, designated S21/MRP, 229,(153) and S21/MRP, 281,(153), encode amino acids 229–1531 and 281–1531 of MRP respectively. The only amino acid introduced during construction of both vectors was an initiator methionine residue. Recombinant bacmids were produced and used to infect Spodoptera frugiperda S21 cells as previously described (Gao et al. 1996).

**Preparation of membrane protein dot blots**

Cells were harvested and membrane-enriched fractions were prepared as described (Hijfner et al. 1994; Gao et al. 1996). Membrane proteins in Tris-buffered saline (TBS) (10 mM Tris–0.15 M sodium chloride, pH 7.5) were blotted onto Immobilon-P polyvinylidene fluoride membrane (PVDF) (Millipore, Mississauga, Ontario, Canada) using a 96-well vacuum manifold. After washing with TBS/0.05% Tween-20 (TBS-T), the blots were cut into strips, transferred to a 24-slot incubation tray and immunoblotted as described below.

**Formic acid hydrolysis of MRP**

Cell membrane-enriched fractions were prepared from MRP-overexpressing H69AR cells. Crude membrane protein (200 μg) was pelleted by centrifugation at 100 000 g and resuspended in 100 μl of 70% formic acid. After incubation at 37 °C for 4 h, the sample was lyophilized and the pellet was washed by resuspension in distilled water followed by lyophilization. The final pellet was resuspended in Tricine sample buffer (BioRad, Hercules, CA, USA) and heated for 45 min at 40 °C. Polypeptides were separated on a 16.5% Tris-Tricine gel, transferred to PVDF and immunoblotted as described below.

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Immunoblotting
Blots were blocked for at least 1 h in 4% skim milk powder in TBS-T, incubated with primary antibody (MAB MRPrl, MAB MRPm6, MAB QCRL-1, MRP-1 or MRP-2 polyclonal antiseras) diluted in blocking solution for 1–18 h and then processed as previously described (Hipfner et al. 1994). Primary antibody binding was visualized by enhanced chemiluminescence detection (Boehringer Mannheim, Laval, Quebec, Canada) using horseradish peroxidase-conjugated F(ab')2 fragments of goat anti-rat IgG (H+L, for MAB MRPrl), goat anti-mouse IgG+IgM (H+L, for MABs MRPm6 and QCRL-1. Pierce, Professional Diagnostics, Edmonton, Alberta, Canada) or goat anti-rabbit IgG (H+L, for antiseras MRP-1 and MRP-2. Jackson ImmunoResearch, West Grove, PA, USA).

Synthesis and immunoblotting of immobilized peptides
Imobilized peptides were synthesized by Research Genetics (Huntsville, AL, USA) using the ‘peptides on paper’ technique. Briefly, peptides were synthesized in nanomolar quantities in a 96-well format, using standard Fmoc chemistry, directly on the surface of an inert membrane. The peptides were covalently linked to the membrane by their COOH-terminal ends via a 6-aminohexanoic acid spacer. For mapping the MAB MRPm6 epitope, 45 decapeptides and one nonapeptide spanning MRP amino acids 1388–1531 and overlapping by seven amino acids [i.e. each successive peptide contained the last seven residues of the preceding peptide plus the next three amino acids in the sequence (or the next two in the case of the nonapeptide)] were synthesized. For mapping the MAb MRPrl epitope, 11 decapeptides spanning MRP amino acids 228–287 and overlapping by five residues were synthesized. The membranes were immunoblotted with MAB MRPm6 or MAB MRPrl essentially as described above, except that 1% bovine serum albumin/2% skim milk powder in TBS-T was used as the blocking solution. MAb MRPm6 and MAB MRPrl were used at concentrations of 2 ng ml⁻¹ and 0.5 ng ml⁻¹ respectively.

Peptide competition of MAB binding
The ability of two peptides (single-letter amino acid code. GSDLWLNKE and PSDLLQQRGL) (Research Genetics) to inhibit binding of MAb to MRP was tested by competitive immunoblotting. Crude membrane proteins (5 μg per lane) prepared from human MRP-transfected HEK cells (HEKGR) (Stride et al. 1997) were separated by SDS-PAGE and transferred to PVDF membrane. The membrane was cut into strips and blocked as above. MAb MRPrl, MRPm6 and QCRL-1 (4 ng ml⁻¹) were mixed with increasing concentrations of peptide GSDLWLNKE (0–200 ng ml⁻¹) or peptide PSDLLQQRGL (0–100 μg ml⁻¹) in 1.5 ml of TBS-T containing 0.1% BSA and 0.005% sodium azide. After incubation for 45 min at room temperature, the MAb/peptide mixtures were added to the strips, incubated overnight at room temperature and the strips processed as above.

RESULTS
Localization of the MAb MRPrl epitope
The approximate location of the MAb MRPrl epitope was determined by immunodot blot analysis of membranes prepared from
Localization of the MAb MRPM6 epitope

We have previously reported that formic acid hydrolysis of MRP is predicted to yield nine fragments ranging in size from 13 to 376 amino acids (Hippner et al. 1996) (Figure 2). The MRP sequence against which MAb MRPM6 was raised (amino acids 1294–1430 and 1497–1531) contains formic acid cleavage sites at amino acids 1376 and 1389. To determine the approximate location of the MAb MRPM6 epitope, we investigated the possibility that its epitope might be located in one of three formic acid fragments of this region. Two of the three fragments are large enough to be resolved by electrophoresis on Tris-Tricine gels. One is a COOH-terminal fragment composed of amino acids 1390–1531 (142 amino acids with a predicted molecular mass of 15.9 kDa) and the other is an internal fragment composed of amino acids 1003–1376 (374 amino acids with a predicted molecular mass of 41.7 kDa). Duplicate blots of MRP-enriched H69AR membranes hydrolysed with formic acid were prepared after SDS-polyacrylamide gel electrophoresis (PAGE) on Tris-Tricine gels. One blot was probed with MRP-2, a polyclonal antiserum that recognizes a sequence (amino acids 1427–1441) predicted to be in the carboxy-terminal formic acid fragment. This antiserum detected a doublet of MRP formic acid fragments of molecular mass 14–16 kDa (Figure 2, left), consistent with the expected sizes of the 142 amino acid carboxy-terminal fragment and a larger 155 amino acid fragment (MRP amino acids 1377–1531) resulting from incomplete hydrolysis. The second blot was probed with MAB MRPM6, which also reacted with two fragments of the same size (Figure 2, right). In view of the fact that the polypeptide used as an immunogen lacked MRP amino acids 1431–1496, these results indicated that the MRPM6 epitope was located between amino acids 1390 and 1430 or between 1497 and 1531.

Epitope mapping with overlapping peptides

Blots of immobilized overlapping peptides encompassing regions of MRP shown to contain the epitopes recognized by MABs MRPr1 and MRPM6 were prepared and probed with the appropriate MAB. Eleven decapeptides overlapping by five amino acids and spanning MRP amino acids 228–287 were synthesized on a membrane and probed with MAB MRPr1. This antibody showed strong reactivity with the peptide GSDLWLSNKE corresponding to amino acids 238–247 of MRP (Figure 3A). Forty-five decapeptides and one nonapeptide overlapping by seven residues and spanning MRP amino acids 1388–1531 were synthesized on a membrane and probed with MAB MRPM6. This antibody showed strong reactivity with the peptide PSDDLQQRGL corresponding to amino acids 1511–1520 near the carboxy terminus of MRP (Figure 3B).

Free forms of the peptides GSDLWLSNKE and PSDDLQQRGL were used in a competitive immunoassay to confirm the specificity of the interactions observed in the blots of immobilized peptides. Immunoblots of membrane proteins from MRP-transfected cells were probed with MABs MRPM6, MRPr1 and QCRL-1 alone or in the presence of increasing concentrations of free peptide. Peptide GSDLWLSNKE inhibited the binding of MAB MRPr1 to MRP in a concentration-dependent manner, and almost completely inhibited binding in this assay at a concentration of 40 ng ml⁻¹ (Figure 4A). In contrast, the binding of MAB QCRL-1 was not affected by this peptide at the highest concentration tested (200 ng ml⁻¹), indicating that the interaction between MAB MRPr1 and Sf21 insect cells that had been infected with recombinant baculovirus expressing two different human MRP constructs with deletions in the region against which this MAB was raised. These deletion constructs, Sf21/MRPós (725–1551) and Sf21/MRPós (751–1551), encode MRP molecules lacking the amino terminal 228 and 280 amino acids respectively. Duplicate blots of membranes prepared from Sf21/MRPós (725–1551) or Sf21/MRPós (751–1551)-infected cells were probed with MAB MRPr1 and MRP-1 polyclonal antiserum. As expected, the MRP-1 antiserum reacted with both the MRPós (725–1551) and MRPós (751–1551) membrane fractions (Figure 1, top) as this antiserum was raised against a peptide in the first NBD of MRP whose sequence is present in both deletion constructs. In contrast, MAB MRPr1 reacted with MRPós (725–1551) membranes but not with MRPós (751–1551) membranes (Figure 1, bottom). These results indicate that the epitope for this MAB was not present in the Sf21/MRPós (751–1551) construct and therefore must be located between amino acids 229 and 280.

and peptide GSDDLWSLNKE is specific. Similarly, peptide PSDLLQQRGL inhibited the binding of MAb MRPrm6 to MRP in a concentration-dependent manner but had no effect on the binding of MAb QCRL-1 at the highest concentration tested (100 μg ml\(^{-1}\)) (Figure 4B), indicating that the interaction between MAB MRPrm6 and peptide PSDLLQQRGL is specific. The locations of the MRPrm6, MRPr1 and QCRL-1 epitopes in the MRP molecule are shown in Fig. 4C.

**DISCUSSION**

MRP is a relatively newly described multidrug resistance protein that, in mammalian cells, confers a phenotype similar to that associated with overexpression of P-glycoprotein, at least with respect to the classes of drugs to which it confers resistance (Loo et al. 1996a; Deeley and Cole, 1997). Consequently, there is considerable interest in determining the relevance of MRP in clinical drug resistance. Several studies examining the expression of MRP in clinical specimens using MRP-reactive MAbs suggest that MRP may be important in a variety of human tumours (Nooter et al. 1995; Filipits et al. 1996a, b; Flens et al. 1996; Chan et al. 1997). In addition to being essential for studies in clinical samples, MRP-reactive MABs have also been extremely useful in structure-function analyses and topological studies of MRP (Bakos et al. 1996; Gao et al. 1996; Hipfner et al., 1996, 1997; Loo et al. 1996a, b, c and 1997). It may be anticipated that, analogous to MABs for other ABC proteins (Shapiro and Ling, 1994; O’Riordan et al. 1995; Illing et al. 1997), they will also prove useful for purification and reconstitution studies of MRP. Given their pivotal role in both clinical and experimental investigations of MRP, it is important that these immunoreagents are well characterized.

With the exception of antisera raised against peptide sequences in the relatively highly conserved NBDs of MRP, none of the MRP-reactive antisera or MABs currently available have been demonstrated to cross-react with P-glycoprotein. This observation is not surprising in view of the very low sequence identity (<15%) between these two proteins. More recently, however, several ABC transporters with substantially greater similarity to MRP than P-glycoprotein have been cloned and characterized. For example, the amino acid sequence of the multispecific organic anion transporter (MOAT) is approximately 50% identical to that of MRP (Paulusma et al. 1996; Taniguchi et al. 1996), and the recently published partial sequences for human MRP3, MRP4 and MRP5 suggest the existence of mammalian ABC proteins with even greater sequence similarity (Kool et al. 1997). Thus, as the ABC superfamily expands and the number of MRP-related proteins increases correspondingly, the specificity of the MRP-reactive MABs becomes less certain. Concerns about cross-reactivity can be greatly alleviated by knowing the epitope sequence recognized by a given MAB, which allows direct sequence comparisons. For example, we have previously determined that the sequence of the QCRL-1 epitope is not found in any other ABC protein characterized to date (Hipfner et al. 1996). Indeed, even though the amino acid sequence of murine mrr is 88% identical to that of human MRP, the QCRL-1 heptapeptide epitope is not conserved in the murine sequence, consistent with the lack of cross-reactivity of this MAB with the murine protein (Hipfner et al. 1996; Stride et al. 1996). Thus, knowledge of the QCRL-1 epitope sequence provides assurance that this MAB is a highly specific probe for human MRP and makes it possible to use free peptide to compete for antibody binding in various immunoassays.

An alignment of the amino acid sequences of the MRPr1 and MRPrm6 epitopes in human MRP with the comparable regions in several MRP-related mammalian ABC proteins shows that, with the exception of murine mrrp, these epitope sequences are poorly conserved (Figure 5). The murine mrrp sequence corresponding to the human MRPr1 epitope differs by only one amino acid, a finding consistent with the reported cross-reactivity of MAB MRPr1 with murine mrrp (Figure 5A) (Flens et al. 1994; Lorico et al. 1996). In view of the low conservation of the MAB MRPr1 epitope in other known MRP-related proteins, it is reasonable to conclude that this MAB is specific for MRP/mrr. The MRPrm6 epitope is also highly conserved in the murine mrrp sequence and differs by only two amino acids (Asp\(^{115}\) → Glu\(^{115}\); Leu\(^{120}\) → Ile\(^{120}\)) (Figure 5B). These differences are very conservative and, consequently, would not necessarily be expected to alter substantially recognition by MAB MRPrm6. Therefore, it is somewhat surprising that this MAB has been reported not to cross-react with murine mrrp, a finding we have confirmed by dot-blot analysis of
mrp-enriched membranes (data not shown). These observations, together with the low level conservation of the epitope sequence among other known MRП-related ABC transporters (Figure 5B), indicate that MAB MRPm6 may be considered a human MRП-specific probe at present.

In conclusion, the data shown in the present study together with those of our previous investigations demonstrate that MABs MRP1, MRPm6 and QCRL-1 recognize highly specific linear epitopes of MRП and, further, that MABs MRPm6 and QCRL-1 are specific probes for the human protein. Because each of the three epitopes is in a different region of MRП, these reagents will continue to be extremely valuable tools for both clinical and structure–function analyses of this protein. We have previously described three additional MABs, namely, MABs QCRL-2, QCRL-3 and QCRL-4, which detect different conformation-dependent intracellular epitopes in the MRП molecule (Hipfner et al. 1994). We have also determined that all three of these MABs are capable of inhibiting the transport function of MRП when measured in inside-out membrane vesicles (Loe et al. 1996b, c, 1997). Accordingly, studies are in progress to map the epitope sequences of these MABs because they are expected to provide important information about the sites and mechanism(s) of substrate binding and transport by MRП.

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REFERENCES

Bakos E, Hegedus T, Hollo Z, Welker E, Tusnady GE, Zaman GJR, Flens MJ, Varadi A and Sarkadi B (1996) Membrane topology and glycosylation of the human multidrug resistance-associated protein. J Biol Chem 271: 12323–12326.

Barnoun K, Leier J, Jedlitschky G, Pourtier-Manzano A, Konig J, Lehmann W-D and Keppeler D (1998) Multidrug resistance protein-mediated transport of chlorambucil and melphalan conjugated to glutathione. Br J Cancer 77: 201–209.

Chan HSL, Lu Y, Grogan TM, Haddad G, Hipfner DR, Cole SPC, Deely RG, Ling V and Galle B (1997) Multidrug resistance protein (MRП) expression in retinoblastoma correlates with rare failure of chemotherapy despite cyclosporine for reversal of P-glycoprotein. Cancer Res 57: 2325–2330.

Cole SPC, Bhardwaj G, Gerlach JH, Mackie JE, Grant CE, Almqvist KC, Stewart AJ, Kurz EU, Duncan AMV and Deely RG (1992) Overexpression of a transporter gene in a multidrug-resistant human lung cancer cell line. Science 258: 1650–1654.

Deely RG and Cole SPC (1997) Multidrug resistance in mammalian cells mediated by members of the ATP-binding cassette superfamily: the P-glycoproteins and MRП. In Molecular Genetics of Drug Resistance, Hasen J and Wolf CR (eds), pp. 247–298. Harwood Academic Press: The Netherlands.

Filipits M, Sachsel RW, Dekan G, Haider K, Valdimarsson G, Depisch D and Parker R (1996a) MRП and MDR1 gene expression in primary breast carcinoma. Clin Cancer Res 2: 1231–1237.

Filipits M, Sachsel RW, Zochbauer S, Malayeri A and Parker R (1996b) Clinical relevance of drug resistance genes in malignant diseases. Leukemia 10: S10–S17.

Flens MJ, Izquierdo MA, Scheffer GL, Fritz JM, Meijer CJLM, Schepers RJ and Zaman GJR (1994) Immunohistochemical detection of the multidrug resistance–associated protein MRП in human multidrug-resistant tumor cells by monoclonal antibodies. Cancer Res 54: 4557–4563.

Flens MJ, Zaman GJR, van der Valk P, Uezuio MA, Schroeijers AB, Scheffer GL, van der Groep P, de Haas M, Meijer CJLM and Schepers RJ (1996) Tissue distribution of the multidrug resistance-associated protein. Am J Pathol 148: 1237–1247.

Gao M, Loe DW, Grant CE, Cole SPC and Deely RG (1996) Reconstitution of ATP-dependent LTC transporter by co-expression of both half-molecules of human MRП in insect cells. J Biol Chem 271: 27782–27787.

Gao M, Yamazaki M, Loe DW, Westlake CJ, Grant CE, Cole SPC and Deely RG (1998) Multidrug resistance protein: identification of regions required for active transport of leuconitriene C4, J Biol Chem 273: 10733–10740.

Giaccone G, van Ark-Otte J, Rubbo GL, Gualdie AF, Brostrom HJ, Dingemans A-M, Flens MJ and Pinedo HM (1996) MRП is frequently expressed in human lung-cancer cell lines, in non-small-cell lung cancer and in normal lung. Int J Cancer 66: 760–767.

Higgins CF (1992) ABC transporters: from microorganisms to man. Ann Rev Cell Biol 8: 67–113.

Hipfner DR, Gualdie SD, Deely RG and Cole SPC (1994) Detection of the M190,000 multidrug resistance protein, MRП, with monoclonal antibodies. Cancer Res 54: 5788–5792.

Hipfner DR, Almqvist KC, Stride BD, Deely RG and Cole SPC (1996) Location of a protease-hypersensitive region in the multidrug resistance protein (MRП) by mapping of the epitope of MRП-specific monoclonal antibody QCRL-1. J Biol Chem 271: 3307–3314.

Hipfner DR, Almqvist KC, Leyle EM, Gerlach JH, Grant CE, Deely RG and Cole SPC (1997) Membrane topology of the multidrug resistance protein, MRП: a study of glycosylation-site mutants reveals an extracellular NH2-terminus. J Biol Chem 272: 23623–23630.

Hilling M, Mokdad LL and Molday RS (1997) The 220-kDa rim protein of retinal rod outer segments is a member of the ABC transporter superfamily. J Biol Chem 272: 10303–10310.

Jedlitschky G, Leier I, Buchholz U, Barnoun K, Kurz G and Keppeler D (1996) Transport of glutathione, glucuronate, and sulfate conjugates by the MRП gene-encoded conjugate export pump. Cancer Res 56: 988–994.

Kool M, de Haas M, Scheffer GL, Schepers RJ, van Eijk MT, Juijn JA, Baas F and Borst P (1997) Analysis of expression of cMOAT (MRП2), MRП3, MRП4, and MRП5, homologues of the multidrug resistance-associated protein (MRП1), in human cancer cell lines. Cancer Res 57: 3537–3547.

Leier I, Jedlitschky G, Buchholz U, Cole SPC, Deely RG and Keppeler D (1994) The MRП gene encodes an ATP-dependent export pump for leuconitriene C4 and structurally related conjugates. J Biol Chem 269: 27807–27810.

Loe DW, Deely RG and Cole SPC (1996a) Biology of drug resistance associated with overexpression of the multidrug resistance protein, MRП. Eur J Cancer 32A: 945–957.

Loe DW, Almqvist KC, Deely RG and Cole SPC (1996b) Multidrug resistance protein (MRП)-mediated transport of leuconitriene C4 and chemotherapeutic agents in membrane vesicles: demonstration of glutathione-dependent vescicle transport. J Biol Chem 271: 9675–9682.

Loe DW, Almqvist KC, Deely RG and Cole SPC (1996c) ATP-dependent 17β-estradiol-17(β-D-glucuronate) transport by multidrug resistance protein (MRП): inhibition by cholesterol steroids. J Biol Chem 271: 9683–9689.

Loe DW, Stewart RK, Massey TE, Deely RG and Cole SPC (1997) ATP-dependent transport of aflatoxin B1 and its glutathione conjugates by the product of the MRП gene. Mol Pharmacol 51: 1034–1041.

Lorico A, Rappa G, Flavell RA and Sartorelli AC (1996) Double knockout of the MRП gene leads to increased drug sensitivity in vitro. Cancer Res 56: 5351–5355.

Mirska SEL, Gerlach JH and Cole SPC (1987) Multidrug resistance in a human small cell lung cancer cell line selected in adriamycin. Cancer Res 47: 2594–2599.

Muller M, Meijer C, Zaman GJR, Borst P, Schepers RJ, Mulder NH, de Vries EGE and Janssen PLM (1994) Overexpression of the gene encoding the multidrug resistance–associated protein results in increased ATP-dependent glutathione S-conjugate transport. Proc Nail Acad Sci USA 91: 13033–13037.

Noordermeer AM, Flens MJ, Zaman GJR, Schepers RJ, van Wingerden KE, Burger H, Oostrom R, Boersma T, Smeenk V, Gratama JW, Kok T, Eggertsson AMM, Bosman FT and Stoter G (1995) Expression of the Multidrug Resistance–Associated Protein (MRП) gene in human cancers. Clin Cancer Res 1: 1301–1310.

Noordermeer AM, Flens MJ, Zaman GJR, Schepers RJ, van Wingerden KE, Burger H, Oostrom R, Boersma T, Smeenk V, Gratama JW, Kok T, Eggertsson AMM, Bosman FT and Stoter G (1995) Expression of the Multidrug Resistance–Associated Protein (MRП) gene in human cancers. Clin Cancer Res 1: 1301–1310.

Oosterluij R. Brucat de la Riviere G, Louk MP. van Wingerden KE.

Henze-Loenhagen HC, Schepers RJ, Flens MJ, Kljin JMG, Stoter G and Foeckens JA (1997) The prognostic significance of expression of the multidrug resistance–associated protein (MRП) in primary breast cancer. Br J Cancer 76: 486–493.

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Norris MD, Bordow SB, Marshall GM, Haber PS, Cohn SL, and Haber M (1996) Expression of the gene for multidrug-resistance-associated protein and outcome in patients with neuroblastoma. *N Engl J Med* 334: 231–238

O'Riordan CR, Erickson A, Bear C, Li C, Manavalan P, Wang KX, Marshall J, Schuske RK, McPherson JM, Cheng SH and Smith AE (1995) Purification and characterization of recombinant cystic fibrosis transmembrane conductance regulator from Chinese hamster ovary and insect cells. *J Biol Chem* 270: 17033–17043

Ota E, Abe Y, Oshika Y, Ozeki Y, Iwasaki M, Inoue H, Yamaazaki H, Ueyama Y, Takagi K, Ogata T, Tamaoki N and Nakamura M (1995) Expression of the multidrug resistance-associated protein (MRP) gene in non-small-cell lung cancer. *Br J Cancer* 72: 550–554

Paulusma CC, Bosma PJ, Zaman GJR, Bakker CTM, Otter M, Scheffer GL, Scheper RJ, Borst P and Oude Elferink RJP (1996) Congenital jaundice in rats with a mutation in a multidrug resistance-associated protein gene. *Science* 271: 1126–1128

Shapiro A and Ling V (1994) ATPase activity of purified and reconstituted P-glycoprotein from Chinese hamster ovary cells. *J Biol Chem* 269: 3745–3754

Stride BD, Valdimarsson G, Gerlach HJ, Wilson GM, Cole SPC and Deeley RG (1996) Structure and expression of the mRNA encoding the murine multidrug resistance protein (MRP), an ATP-binding cassette transporter. *Mol Pharmacol* 49: 962–971

Stride BD, Grant CE, Loe DW, Hipfner DR, Cole SPC and Deeley RG (1997) Pharmacological characterization of the murine and human orthologs of multidrug resistance protein (MRP) in transfected human embryonic kidney cells. *Mol Pharmacol* 52: 344–353

Taniguchi K, Wada M, Kohno K, Nakamura T, Kawabe T, Kawakami M, Kagotani K, Okamura K, Akiyama S and Kuwano M (1996) A human canalicular multispecific organic anion transporter (cMOAT) gene is overexpressed in cisplatin-resistant human cancer cell lines with decreased drug accumulation. *Cancer Res* 56: 4124–4129