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Temporal Profiling of the Coding and Noncoding Murine Cytomegalovirus Transcriptomes[⊽]†

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The global transcriptional program of murine cytomegalovirus (MCMV), involving coding, noncoding, and antisense transcription, remains unknown. Here we report an oligonucleotide custom microarray platform capable of measuring both coding and noncoding transcription on a genome-wide scale. By profiling MCMV wild-type and immediate-early mutant strains in fibroblasts, we found rapid activation of the transcriptome by 6.5 h postinfection, with absolute dependency on *ie3*, but not *ie1* or *ie2*, for genomic programming of viral gene expression. Evidence is also presented to show, for the first time, genome-wide noncoding and bidirectional transcription at late stages of MCMV infection.

Murine cytomegalovirus (MCMV) is a ubiquitous betaherpesvirus with a 235-kbp genome transcribed in a classical cascade fashion (55). The genome sequence of MCMV has been available for some time (89), and yet a systematic study of temporal gene expression during MCMV infection has been lacking. Double-stranded PCR-based cDNA microarrays were used once previously (105) to validate the expression of a subset of predicted MCMV open reading frames (ORFs) (15) at a single time point (24 h postinfection [hpi]). More-advanced microarray technology based on oligonucleotide probes, affording increased specificity to distinguish RNA polarity, has not been reported. For human cytomegalovirus (HCMV), abundant antisense (AS) transcription has been observed (120), raising the possibility that aspects of the CMV life cycle are influenced or regulated by noncoding transcripts. For MCMV, small virus-encoded microRNAs (miRNAs) (19, 20, 33) and larger double-stranded RNAs (21, 110) have been reported to be transcribed from multiple loci; however, the frequency and abundance of noncoding transcripts throughout the MCMV genome have not yet been measured systematically on a genome-wide scale and at multiple stages of infection.

Here we have investigated the global transcriptional program of MCMV by constructing a microarray capable of measuring sense (S) and AS transcripts. Microarrays were designed using 55-mer oligonucleotide probes in sense and antisense orientations to each of the 170 viral ORFs predicted in the MCMV genome (89). One hundred ninety-two positive-control probes were designed against stably expressed mouse genes for normalization purposes, and 97 negative-control probes were designed against Saccharomyces cerevisiae sequences with no homology to mouse or MCMV genomes (for probe sequences, see Table S1 in the supplemental material). The 55-mer oligonucleotide probes were diluted to a concentration of 60 µM and inkjet printed (Arrayjet, United Kingdom) onto amino silane-coated glass slides, with each microarray consisting of six identical subarrays. Probes were printed in triplicate per array and have the capacity for developing a total of 18 measurements per probe per sample to ensure high technical replication. Target RNA was extracted from infected fibroblasts using PureLink RNA minikits with on-column DNase I treatment (Invitrogen, CA). Purified RNA (700 ng for each sample) was labeled for microarray analysis using the Agilent low-input fluorescent linear amplification protocol (Agilent, CA), with 3 µg of Cy5-labeled target cRNA hybridized per sample. Hybridized microarrays were washed and subsequently scanned using an Agilent (CA) G2505B scanner.

To perform a systematic analysis of genome-wide transcription in MCMV, we infected NIH 3T3 fibroblasts with the parental MCMV strain at a multiplicity of infection (MOI) of 1 and performed DNA microarray analysis on total RNA harvested from duplicate cultures at 0.5, 6.5, 24, and 48 hpi. Individual probe signals were background subtracted, median summarized, and log base 2 transformed to form raw data points (see Table S2 in the supplemental material). Raw data were quality controlled, and normalization between samples

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sense probes ^a
microarray
MCMV
High-confidence
1.
TABLE

	Virion association ^e								VAP						VAP	VAP	VAP VAP	VAP VAP	VAP	VAP VAP	VAP	VAP	
roarray sense propes	Annotation ^d (reference[s])	Spliced m128 (ie2) gene has sequence similarity to members of the US22 family of HCMV (77)	ie1 exon 4, with mRNA terminating at base 179544 (53-55); total length of Ex2 plus Ex3 plus Ex4 is 595 aa, total molecular size is 66.7 kDa	Sequence variation and early transcriptional kinetics found in wild-type-derived MCMV isolates: m03-encoded protein could be found on cell surface (28)	m04 gene product (gp34) forms a complex with MHC-I (51), which reaches the cell surface (46) and is required for Ly49P recognition of infected cells (57); m04(m34 also antaomizes the effect of m173 (84)	motive provides the provident of must contract of MHC-1 proteins in motive product (grays) downmodulates the levels of MHC-1 proteins in infected cells (50) and binds to the ternary complex of assembled MHC-1 with antigenic peptide in the ER, reduce thing this complex of the lysosome for dominant complex of an end of a dominant of the second of the s	degradation; unbound gp48 is destroyed by the proteasome (18) Glycoprotein family $m02$	Glycoprotein family m02 Glycoprotein family m02, probe overlaps with newly predicted ORF m16.2 (105)	Highly antigenic early gene (44)	Mutant showed no obvious growth phenotype (75)	Reannotations of the MCMV genome have identified three putative M34- overlapping ORFs (m331, m341, and m34.2); this microarray probe overlaps with newly predicted ORF m34.2 (105); an M34 mutant virus which interrupted all three m34 ORFs had attenuated replication both in tissue	culture and m SCLD mice (10) Possible transcriptional regulator (89), implicated in blocking apoptosis via inhibition of caspase-8 (FLICE) activation (100); growth of M36 mutant was attenuated in vitro and in vivo (27); HCMV VICA/pUL36 protects cells from	Gycoprotein, wIIA: $M37$ mutant is severely attenuated in growth and virulence of ycoprotein, wIIA: $M37$ mutant is severely attenuated in growth and virulence in vivo (63); homolog of HCMV UL37 that inhibits mitochondrial megapore activation in a manner similar to members of the antiapoptotic Bcl family	(37); may also be a transcriptional regulator (58) Putative antiapoptotic function (79)	Antiapoptotic (79), immunoregulatory gene that modulates T helper cell response (99); found to be nonessential for viral growth <i>in vitro</i> and <i>in vivo</i>	and dispensable for virulence in killing SCID mice (117) Antiapoptotic (17) homologue of the large subunit of ribonucleotide reductase (85); blocks NF-kB activation as a result of its inhibitory effect on RIP1 (55); m_{1000}	sugarang (11, 109) Smallest capate protein (14) DNA polymerase delta subtype (59)	Major ssDNA binding protein (3, 4) Assembly protein and protease (11, 60) that conserves the domain structure and chomono citics account in UCMV1 II 60.	the ways are a present in TICMY CLOOD Encodes a structural protein unique to the betaherpesvirus group, also known as	Vip) 1, same taumy as MO2 (21) Vipor-associated, tegument/second envelopment protein (112) Encodes small structural phosphoprotein unique to the betaherpesvirus group, shown to be around 16 kDa in size and to be associated with the virion (30,	78) Part of the helicase-primase complex of three proteins $(M70, M102, and M105)$	(M, M) but A helicase; part of the helicase-primase complex of three proteins (M70, M, M, M	Possible alternate splice to m119, as for HCMV UL118 (62, 88) Exon 2 of M133 (sgg1) (61, 73)
ign-confidence incluiv mic) Protein type	Immediate early	Immediate early	Glycoprotein m02	Glycoprotein m02	Glycoprotein m02	Glycoprotein m02	Glycoprotein m02 Glycoprotein m02	Membrane glycoprotein Early	US22 family homolog		US22 family homolog	Glycoprotein	Putative eleconrotein	US22 family homolog	Ribonucleotide reductase	Capsid DNA polymerase	DNA binding Assembly protein, protease	Upper matrix	Virion associated Phosphoprotein	Helicase-primase	DNA helicase	
ABLE I. N	Time on ^c (hpi	0.5	6.5	6.5	6.5	6.5	6.5	6.5 6.5	6.5 6.5	6.5 6.5	6.5	6.5	6.5	6.5 6.5	6.5	6.5	6.5 6.5	6.5 6.5	6.5	6.5 6.5	6.5	6.5	6.5 6.5 6.5
Τ	e HCMV name	US22 (GF2)	IE1							UL23 (GF2) UL28	UL34	UL36 exon 2	NT37		UL43	UL45 (RRL)	UL54 (DNApol)	UL57 (MDBP) UL80 (AP)	UL82 (pp71)	UL94 UL99 (pp28)	UL102 (HP)	UL105 (Hel)	UL118
	MCMV name	ie2	ie1		gp34	gp48											DNApol		pp71				sgg1
	Strand	C	U					C	ວບ	υu		C	C	00	U C	C	υu	U	C				000
	ORF 5	m128 Ex3	m123 Ex4	m03	m04	m06	m08	m10 m16.2	m17 m18	m25.1 M28	m34.2	M36 Ex2	M37	m41 m42	M43	M45	m48.2 M54	M57 M80	M82	M94 M99	M102	M105	M118 m119.2 m132 Ex2
	Unique probe ID ¹	vMC132	vIE1rem	vMC003	vMC004	vMC006	vMC008	vMC010 vMC016	vMC017 vMC018	vMC027 vMC030	vMC037	vMC040	vMC041	vMC045 vMC046	vMC047	vMC049	vMC055 vMC061	vMC064 vMC079	vMC080	vMC092 vMC098	vMC100	vMC103	vMC114 vMC117 vMC136

owing page	ICP18.5 (13, 38) Continued on fol	2		~			
VAP VAP	Conserved herpesvirus protein that forms a complex essential for egress of nucleocapsids from the nucleus (94); M50/p35 recruits cellular PKC for nucleocapsids from the nucleus (94); M50/p35 recruits cellular PKC for Glycoprotein B (29, 87) Tegument protein and homologue of HCMV terminase subunit and HSV	Nuclear export 0 Glycoprotein 0 Tegument 1	24 24 24		UL53 UL55 (gB) UL56 (NM)	BB UL53 UL55 (gB) UL56 (NM)	UL53 C gB UL55 (gB) C gB UL56 (NM)
	membrane and is aggregated by $M53/p38$ to form the capsid docking site (82)		24		UL52	UL52	NL52
	Conserved herpesvirus protein that forms a complex essential for egress of mucleaverasids from the nuclear (94): M50/h55 inserts into the inner nuclear	Nuclear export	24		UL50	ÚL50	C UL50
VAP	Large tegument protein (101)	Tegument	1 4 K		UL48 (Teg) 111 40	UL48 (Teg) 111 A0	UL48 (Teg)
	US22 family homolog	1	4 4 6 4 4 6		UL36 exon 1 UL38	UL36 exon 1 UL38	UL36 exon 1 C UL38 exon 1
IAJ	Virulence factor and HCMV UL25 family homologue (26, 32); M35 insertional	Virion associated	24		UL35 (GF1)	UL35 (GF1)	UL35 (GF1)
Ň	errors in the 5 -proximat end (+9) Encodes large phosphoprotein homologous to HCMV tegument protein UL32 (150) (177–80)	Phosphoprotein (tegument) H	24		UL32 (pp150)	UL32 (pp150)	C UL32 (pp150)
	Reported irregularities in coding potential have suggested possible sequence	ł	244 244		UL31	UL31	C UL31
	Plays important role in MCMV growth and virulence (2) and inhibits IFN-y sionalino (56) via selective hinding and downroutlation of Stat? (133)	H	24 24		UL26 UL27	UL26 UL27	C UL26 C UL27
	UL25 family homolog UL25 family homolog (110)	Tegument L	24 24		UL23 (GF2) P UL25 (GF1)	UL23 (GF2) P UL25 (GF1)	C UL23 (GF2) P UL25 (GF1)
	Reported irregularities in coding potential have suggested possible sequence	Glycoprotein m02 Glycoprotein m02 Glycoprotein m02 F	2 2 2 4 2 4 4 4				U
	ie3 exon 5, with mRNA terminating at base 177817 (76); total length of Ex2 plus Ex3 plus Ex5 is 611 aa. molecular size is 68.1 kDa	Immediate early	6.5 24		UL122 (IE2)	ie3 UL122 (IE2)	C ie3 UL122 (IE2)
	1-265 is presented by the MHC-1 molecule D(d) (45) Required for MCMV virulence in killing SCID mice and for optimal viral	glycoprotein Putative membrane	6.5				C
	MCMV viral carrier protein gp36.5 (64); m164-derived peptide 257-AGPRYSR	glycoprotein Putative membrane	6.5			gp36.5	C gp36.5
		Putative membrane glycoprotein Putative membrane	c.0 2.0				0 0
	Aming in mice (+2) Threonine-serine-rich glycoprotein of MGP family m145, some homology to EHV1 g(N) (106)	Glycoprotein	6.5				C
	MHC-I proteins in infected cells (50), disrupts export of MHC-I complexes from pre-Golgi compartments to the Golgi compartment via a luminal domain (122), and modulates NK cell response (inhibition), antigen presentation, and T cell response (40); viruses lacking gp40 show increased susceptibility to CTL Fillion in <i>inco</i> (43)						
-	Spliced gene m147.5 selectively targets CD86 expression on APCs (68)	Possible membrane-spanning Sprotein	6.5				C
	deletion mutants show unexpected alterations in virulence and are attenuated in normal and immunosuppressed adult mice (31); m138 downmodulates the NKG2D ligands MULT-1, H60 (65), and RAE-1 epsilon (8) Required to block PKR-mediated shuldown of cellular protein synthesis and	US22 family homolog	6.5		US26 (GF2)	US26 (GF2)	C US26 (GF2)
	Encodes the 88-kDa Fc receptor glycoprotein (107); Fc receptor-specific (m138)	Putative glycoprotein Glycoprotein	6.5 6.5		fcr1	Fcgr fcr1	C Fcgr fcr1

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1-Cor	
LABLE	

	Virion association ^{e}	VAP	VAP		VAP	VAP VAP VAP	VAP	VAP VAP												
	Annotation ^d (reference[s])	Tegument protein similar to HCMV transactivator UL69 (113), which in HCMV	induces a G ₁ block (42) Honology to HCMV UL72, a putative dUTPase enzyme required for nucleotide metabolism replication 20, a nad/or repair (86)	G protein-coupled receptor homologue, same family as M33 (38); has subcellular trafficking properties (96); M78 mutants exhibit reduced replication in cultured	Homologue of HCMV virion-associated factor with IFN repressor function (pp65) (1, 16, 74); M83 mutant has attenuated viral growth and virulence in	SCID mice (119) Homologue of HCMV minor capsid protein (9) Homologue of HCMV major capsid protein (25) Homologue of HCMV virion protein (9)	Homologue of HCMV UL97 phosphotransferase gene, whose product	phosphorylates ganciclovir in HCMV-infected cells (67, 102) Alkaline exonuclease (DNase) gene (89) Glycoprotein M with seven hydrophobic stretches that are potential membrane-	spanning regions (66, 93)	Exon 1 of e1 (22); total e1 length is 330 as, molecular size is 36.4 kDa; IE3 and	the early M11.2/115 gene products colocatize and communoprecipitate (1/94), neuron-specific activation of the e1 promoter observed in transgenic mice (7) Uracil DNA glycosylase enzyme homolog (114) found in herpesviruses, required	for nucleotide metabolism, replication, and/or repair M115 (gL) (52) contains five potential glycosylation sites, has significant amino acid similarity to gL homologs in HCMV and HHV-6, and has previously been shown to he advoscitated in virians (118)	Predicted MCMV homologue of HCMV spliced ORF (89)	ie3 exon 5, with mRNA terminating at base 177817 (76); total length of Ex2 plus Ex3 alone Ex5 in 611 nor molecular give is 68 1 bDo	Explore EX is our ad, inorecutal size is our taxa ORFs m124, m124.1, and m125 located within the enhancer region are nonessential for MCMV growth <i>in vitro</i> (6)	Exon 2 of M133 (sgg1) (61, 73) US22 family homolog Downmodulates expression of cellular MULT-1 (60) and is a member of MGP	Tamuy m145 Member of MGP family m145 that modulates NK cell (inhibition) and T cell	response and impedes an NKG2D ligand (H60) (41, 79)	Putative membrane glycoprotein	
TABLE 1—Continue	Protein type	egument	utative dUTPase	protein-coupled receptor	irion associated	apsid apsid irion associated	hosphotransferase	xonuclease lycoprotein		arly	lycosylase	lycoprotein	otential glycoprotein	nmediate early		lycoprotein	lycoprotein	utative membrane	glycoprotem utative glycoprotein	lycoprotein m02 lycoprotein m02 lycoprotein m02
	lime on ^e (hpi)	24 T	24 Pı	24 24 G	24 24 V	400 7444 400	244 244 P	24 24 G	24 24 24	24 24 E	24 G	24 G	22224 4444 72244 7	24 Ir	24	24 24 24 G	24 24 G	24 P1	24 24 Pi	48 48 48 48 48 48 48 48 48 48 48 48 48 4
	HCMV name T	JL69	JL72 (dUTPase)	JL73 JL78	JL79 JL83 (pp65)	JL85 JL86 (MCP) JL88 J103	11.89 11.95 11.97 (PK)	JL98 (DNase) JL100 (gM)	JL103	JL112	JL114 (UNG)	JL115 (gL)		JL122 (IE2)		JS22 (GF2)				
	MCMV name		L					g Mg	Ļ	el L		gL L		ie3 L		sgg1 L				
	Strand]	C	C		υu	υu	U	C	00	C	C	С	C	JU		υu		C	C	C
	ORF	M69	M72	M73 M78	M79 M83	M85 M86 M88 M03	M95 M95 M97 M97	M98 M100	M103 m106 m107	m108 M112 Ex1	M114	M115	m119 m119.1 m119.4 m119.5	M122 Ex5	m124	m132 Ex1 m139 m145	m148 m155	m159	m161 m168	m170 m15 m12 m14
	Unique probe ID^b	vMC067	vMC071	vMC072 vMC077	vMC078 vMC081	vMC083 vMC084 vMC086	vMC093 vMC094 vMC095 vMC096	vMC097 vMC099	vMC101 vMC104 vMC105	vMC106 vMC107	vMC109	vMC110	vMCI15 vMCI16 vMCI19 vMCI20	vMC123	vMC127	vMC137 vMC143 vMC149	vMC152 vMC159	vMC163	vMC165 vMC172	vMC174 vMC005 vMC012 vMC012

VAP			nded DNA;
Potential ORF located within the origin of replication (89) Functional homolog of HCMV gO, has key role in determining the entry	pathway of MCMV (95) Potential alternative splice to M112 Ex1 (e1) (22), as found for HCMV UL112/ TUT112 (1157)	Predicted MCMV homologue of HCMV spliced ORF m131/129 is a chemokine homolog and determinant of viral pathogenicity (35) and may modulate cytokine signaling (79)	mock-infected cells to a high confidence level ($P \le 0.05$ by empirical Bayes analysis). endoplasmic reticulum; RIP1, receptor-interacting protein kinase 1; ssDNA, single-stra IFN- γ , gamma interferon; HHV-6, human herpesvirus 6.
Putative glycoprotein			in MCMV-infected cells versus rrial inhibitor of apoptosis; ER, ; CTL, cytotoxic T lymphocyte;
4 4 4 8 8 8	48	48 8	pregulated I mitochonc
UL74 P	UL113 P		RFs found to be significantly u aspase-8 activation; vMIA, vira nting cells; MGP, membrane g l on reference 49.
υυ		C	ng MCMV O inhibitor of c antigen-prese (VAPs), basec
m39 m59 m74	M113	m119.3 m129	obes for codii be identifier. i detected. Is; vICA, viral ase R; APCs, ted proteins (
vMC043 vMC066 vMC073	vMC108	vMC118 vMC133	^a Microarray pr ^b Probe ID, pro ^c Time on, time ^d aa, amino acic PKR, protein kim ^e Virion-associa

was performed based on a subset of 44 positive-control probes highly correlated across the data set (Pearson r of >0.90). Normalized expression data (see Table S3 in the supplemental material) were subjected to a statistically rigorous threshold detection methodology for providing on/off calls for each probe based on a receiver operating characteristic (ROC) (12). From these ROC analyses, we evaluated specificity levels corresponding to given sensitivities of 70%, 80%, and 90%. At a moderate sensitivity of 70%, we were able to obtain an average specificity of 93%, and this was chosen as affording an optimal balance between identifying true positives and excluding true negatives with stringency (for ROC plots, see Fig. S1 and S2 in the supplemental material). Accordingly, we detected 297 total probes having "on" calls and 163 probes for coding MCMV ORFs, making 87.6% of the MCMV genome detectable at 48 hpi (for a list of genes detected, see Table S4 in the supplemental material).

To account for experimental variation, statistical testing (empirical Bayes moderated t test) was applied between mockinfected and infected groups to identify differential expression of only the most highly significant MCMV ORFs. By use of this more stringent approach, 119 ORFs were found to be significantly activated to a confidence level of $P \le 0.05$ above mockinfected levels at all time points (Table 1). These included the DNA polymerase subunit M54 (59), known inhibitors of major histocompatibility complex class I (MHC-I) surface expression m04 (gp34) (51) and m06 (gp48) (90), and the Fc receptor m138 (108). After a single round of replication at 24 hpi, a total of 111 MCMV ORFs were detected at the high significance level. To further validate these findings, a subset of MCMV ORFs were subjected to quantitative reverse transcription-PCR (qRT-PCR) analysis (for primer sequences, see File S1 in the supplemental material), and in agreement with the microarray results, each test case showed that ORF expression was also detectable by qRT-PCR (Fig. 1a).

As previously noted for HCMV microarray analysis, there is no overt positional bias toward expression of coding ORFs based on genomic location that could be linked to the patterns of gene expression observed during infection. MCMV ORFs were annotated based on Rawlinson et al. (89) and updated with details from additional publications wherever possible. MCMV ORFs from recent predictions (15, 105) were aligned against 55-mer probe sequences, identifying five probes overlapping with newly predicted ORFs (m107-m107.2, m16m16.2, m22-m22.1, M34-m34.2, and M58-m58.1), which were reannotated accordingly.

As a result of the statistical cutoff ($P \le 0.05$), MCMV probes for ORFs M44, M70, M75, m135, m143, m144, m153, and m157 failed to be included, although these genes have been reported to be expressed in previous MCMV studies or are homologues of HCMV genes reported to be expressed (21, 29, 34, 79, 80, 89, 92, 110, 118). It is most likely that the specific probes for these genes exhibit false-negative results. Nevertheless, in this study, we aimed to purposefully avoid false positives at the sacrifice of capturing a modest level of false negatives. For this reason, we also did not detect iel or ie3 expression until 24 hpi based on the statistical cutoff; however, these genes are detectable as early as 0.5 hpi and 2.5 hpi by use of a more sensitive qRT-PCR approach (see Fig. S3 in the supplemental material).



FIG. 1. MCMV ORF detection and characterization of MCMV IE deletion mutant strains. (a) (Panel i) Microarray signals for 10 MCMV ORFs in mock- and MCMV-infected cells. (Panel ii) qRT-PCR validation of the 10 MCMV ORFs at 24 hpi. The *y* axis represents crossing point (Cp) values from qPCR amplification curves, with low Cp values indicating high transcript abundance. Error bars represent standard errors of the mean. (b) Schematic representations of recombinant MCMV IE deletion mutants. The map of the parental MCMV genome is shown at the top, with structures of the *ie1*, *ie2*, and *ie3* transcripts below. Coding exons are shown in black, with arrows indicating the directions of transcription, and white triangles represent deleted loci. The gray box marks the MCMV *ie1/ie3* promoter enhancer (MIEP). (Line iii) The MCMVdie3 revertant strain was renamed MCMVdiie2 in this study, as the HCMV MIEP is inserted between two HpaI sites spanning the transcription start site of the *ie2* gene, disrupting *ie2* expression (marked with ×). GFP, green fluorescent protein. (c) Viral titers and genome particle/PFU equivalences of the four MCMV strains, as determined by qPCR.

To gain further insight into the transcriptional programming of MCMV, we next sought to profile gene expression from three well-characterized MCMV mutants (5, 23, 36, 72) alongside the parental MCMV strain (111) (for schematics of strains, see Fig. 1b). To characterize the mutant strains before microarray analysis, we sought to (i) determine equivalent infectious doses at the genomic level by measuring genome/PFU ratio for each stock, (ii) ensure that generating the *ie3* deletion mutant (MCMVdie3) in a complementing cell line did not drastically alter the infectious-particle ratio, (iii) ensure that viral growth phenotypes were consistent with those previously published (5, 36, 72), and (iv) ensure that no viral transcription was occurring from deleted loci. On the basis of quantitative PCR (qPCR), we detected equivalent numbers of MCMV genome copies per PFU for each viral strain (Fig. 1c), using as a calibrator a reference plasmid containing the m115 (gL) gene (nucleotides [nt] 166387 to 167208; GenBank accession no. NC 004065) (for a detailed account of this approach, see reference 98). Equivalent numbers of MCMV genomes were also found at 2 hpi inside cells infected with different MCMV strains (data not shown). Multistep growth curves confirmed viral growth phenotypes (see Fig. S4 in the supplemental material), and qRT-PCR (Qiagen 1-step kit; Germany) confirmed that transcription was not detectable from the respective deleted loci for *ie1*, *ie2*, and *ie3* (data not shown). These data show that the four MCMV strains are experimentally comparable for downstream gene expression analysis.

Based on previous virologic characterization of these MCMV immediate-early (IE) mutants, we expected to observe different gene expression profiles for the MCMV strains. For example, given that removal of the ie2 gene causes no reported change in phenotype (45), we anticipated few gene expression changes in the MCMVdiie2 strain relative to the parental MCMV strain. Alternatively, given that ie3 has an indispensable regulatory function and is essential for viral growth (5), we expected little or no viral gene expression to be detectable from the MCMVdie3 strain. However, for the ie1 deletion mutant (MCMVdie1), the expectation was less clear, given that this strain has wild-type growth characteristics and yet the IE1 protein is well known to have transcriptional regulatory activity (24, 39, 76) and is further known to interact with cellular host factors (81, 104). In order to profile the gene expression of each mutant strain, we infected NIH 3T3 fibroblasts in parallel at an MOI of 1 and harvested total RNA for microarray anal-



FIG. 2. Gene expression program of MCMV, MCMVdie1, MCMVdiie2, and MCMVdie3. (a) Volcano plots comparing microarray signals from mock-infected and infected samples at 48 hpi, using *P* value (*y* axis) and fold change (FC) (*x* axis) comparisons derived from empirical Bayes testing, with two biological replicates per group. (b) Hierarchical clustering of high-confidence MCMV probes, with each row representing a single probe normalized to its mean value across the data set to show relative expression. Yellow indicates increased expression and blue indicates decreased expression relative to the mean. Genes clustered based on the similarity of their expression profiles across the data set, with similar genes connected at the hierarchical tree on the left. Data represent mean values from two biological replicates. Numerical values at top indicate hours postinfection.

ysis at 0.5, 6.5, and 48 hpi, along with that of mock-infected cells.

Figure 2 shows the comparative activation of viral transcriptomes among the four MCMV strains at 48 hpi and indicates that the transcriptomes of MCMVdie1 and MCMVdiie2 are activated with profiles of viral gene expression very similar to that of the parental MCMV strain over the 48-h period (Fig. 2a). Similar numbers (within 10%) of MCMV probes were detectable from MCMVdie1 (103 ORFs), MCMVdiie2 (114 ORFs), and the parental strain (113 ORFs) at 48 hpi. The degree of similarity in expression profiles among the MCMVdie1, MCMVdiie2, and parental MCMV strains suggests that *ie1* and *ie2* have a redundant or negligible transcriptional regulatory role in controlling downstream MCMV gene expression during fibroblast infection. Hierarchical clustering (Fig. 2b) further indicates few, if any, differences in the global gene expression profiles of MCMVdie1 and MCMVdiie2 compared to that of the parental MCMV strain. These results could point to a redundant role for the *ie1* and *ie2* genes in controlling downstream viral gene expression or, alternatively, a lack of

TABLE 2. High-confidence MCMV	microarray antisense probes ^a
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Unique probe ID ^b	ORF	Time on ^c (hpi)	Details of cDNA cloning validation	Sense probe ^d
vMC336	m148as	6.5	Validated with AS cDNA clone L151 at nt 208012 to 207467 (5' to 3'), overlapping m148 (AS) and m149 (S); clone length, 545 nt	S
vMC357	m169as	6.5		S
vMC175	m05as	24		S
vMC180	m163as	24		S
vMC188	m04as	24	Validated with AS cDNA clone IE150 at nt 4043 to 3943 (5' to 3'), overlapping m04 (AS); clone length, 384 nt	S
vMC189	m05as	24		S
vMC190	m06as	24		S
vMC191	m07as	24		
vMC193	m09as	24		
vMC197	m13as	24		S
vMC229	m41as	24		S
vMC236	M47as	24		
vMC241	M50as	24		S
vMC248	M57as	24		S
vMC257	m74as	24	Validated with AS cDNA clone L147 at nt 104825 to 105449 (5' to 3'), overlapping m74 (AS); clone length, 624 nt	S
vMC269	M87as	24		
vMC270	M88as	24		S
vMC283	M102as	24		S
vMC285	M104as	24		
vMC291	M113as	24		S
vMC293	M115as	24		S
vMC299	m119.2as	24		S
vMC311	m124as	24		S
vMC333	m145as	24		S
vMC335	m147as	24		
vMC351	m163as	24		S
vMC356	m168as	24		S
vMC239	M48as	48		S
vMC251	m69.1as	48		
vMC272	M89 Ex2as	48		
vMC277	M94as	48	Validated with AS cDNA clone L164 at nt 137299 to 137096 (5' to 3'), overlapping m94 (AS); clone length, 203 nt	S
vMC289	m108as	48		S
vMC300	m119.3as	48		S
vMC319	m132 Ex2as	48		S
vMC332	m144	48		

^{*a*} Microarray probes for antisense transcripts found to be significantly upregulated in MCMV-infected cells versus mock-infected cells to a high confidence level ($P \le 0.05$ by empirical Bayes analysis).

^b Probe ID, probe identifier.

^c Time on, time detected.

^d AS probes with significant signal also found from the corresponding sense probes are marked with an "S."

sensitivity in controlling the viral genomic program in fully permissive fibroblasts.

In marked contrast to the *ie1* and *ie2* mutant strains, MCMVdie3 exhibited an undetectable level of viral gene expression, suggesting that *ie3* acts as a global *trans*-activator of downstream MCMV gene expression, as indicated by previous studies (5). To further examine the transcriptional status of MCMVdie3 using a more sensitive approach, IE and downstream MCMV genes were measured using qRT-PCR in MCMVdie3-infected cells both in the presence and in the absence of 50 μ g/ml cycloheximide (C7698; Sigma, United Kingdom) at 2.5 hpi. These experiments confirmed that IE kinetic class genes were expressed in MCMVdie3 but that genes beyond the IE region were not (see Fig. S5 in the supplemental material).

The design of the MCMV microarray platform enables selective detection of transcripts originating from both strands of the viral genome by having probes designed in sense (S) and antisense (AS) orientations to each MCMV ORF. At 24 hpi, we detected antisense transcripts from 23 AS loci, five of which (m104as, M113as, m147as, m163as, and m168as) have overlapping ORFs on the opposite strand of the genome, indicating known or predicted regions of bidirectional transcription based on prior annotation (105). Three other loci were found to have neighboring but nonoverlapping ORFs in their vicinity (M57as, m74as, and M88as). An additional 15 MCMV AS probes detected at 24 hpi were found to have no overlapping or nearby ORFs located on the opposite strand, indicating previously unknown noncoding transcripts derived from regions outside MCMV ORFs (m04as, m05as, m06as, m07as, m09as, m13as, m41as, M47as, M50as, M87as, M102as, M115as, m119.2as, m124as, and m145as). At 48 hpi, an additional eight antisense probes were detectable, but all have overlapping or nearby ORFs on the opposite strand of the genome (M48as, m69.1as, M89as, M94as, m108as, m119.3as, m132as, and m144as). In total, evidence of antisense transcription was detected from 35 loci over the four time points as measured by microarray analysis (Table 2 and Fig. 3). Twenty-six of these loci were also



FIG. 3. MCMV genome activation measured by microarray analysis and qRT-PCR. Transcript abundance of ORFs expressed from the parental MCMV strain was measured using oligonucleotide microarrays at 0.5, 6.5, 24, and 48 hpi in NIH 3T3 cells at an MOI of 1. Histograms represent mean values from two replicate samples after background (mock infection) subtraction. Transcripts are arranged in order from left to right according to ORF names ranging from m01 to m170, with sense probes shown on the left and antisense probes on the right. All raw data are available in the supplemental material.

found to have significant signal from the corresponding sense (S) probe, indicating a potential site of bidirectional transcription. A trend toward antisense transcription occurring more frequently at the terminal ends of the MCMV genome is also noted (Fig. 3).

In order to independently validate AS transcripts identified by microarray analysis, we generated cDNA libraries from MCMV-infected fibroblasts pooled from 4, 8, and 12 hpi (IE library), 16, 24, and 32 hpi (E library), or 40, 60, 80, and 100 hpi (L library). cDNA libraries were generated as described previously for HCMV (120). cDNA clones overlapping AS microarray regions were found at m04as, m74as, M94as (none of which have overlapping ORFs on the opposite strand of the genome), and m148as (which has the m147 ORF on the opposite strand of the genome) (for validated AS transcripts, see Table 2). We also found one large cDNA clone that overlapped three MCMV ORFs (m119.2, m119.3, and m119.4), two of which had potential AS regions identified by microarray probes (m119.2 and m119.3). Four additional AS cDNA clones were found to overlap AS regions not identified by microarray analysis (m19as, M72as, m149as, and m151as). These experiments thus reveal for the first time that antisense transcription occurs frequently throughout the MCMV genome, an observation that will likely seed further studies.

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