

TRANSACTION GRANTED EARLY TERMINATION—Continued

ET date	Trans No.	ET req status	Party name	
28-SEP-09	20090747	G	International Business Machines Corporation.	
		G	Court Square Capital Partners II, L.P.	
		G	Rocket Software, Inc.	
	20090757	G	G	Rocket Software, Inc.
			G	H.I.G. Capital Partners IV, L.P.
			G	Stant Parent Corp.
		G	G	Standard-Thomson Corporation.
			G	Stant Holding Corp.
			G	Thomson International Corporation.
	20090770	G	G	Stant Manufacturing, Inc.
			G	Stant Corporation.
			G	M. Brooks Smith.
20090771	G	G	Ami Shashoua.	
		G	QPay, Inc.	
		G	M. Brooks Smith.	
29-SEP-09	20090742	G	Yossi Amosy.	
		G	QPay, Inc.	
		G	Hon Hai Precision Industry Co., Ltd.	
30-SEP-09	20090732	G	Sony Corporation.	
		G	Sony Electronics Inc.	
		G	Sony Baja California, SA. de C.V.	
		G	Warner Chilcott plc.	
		G	The Procter & Gamble Company.	
		G	Procter & Gamble Pharmaceuticals Longjumeau SAS.	
		G	Procter & Gamble S.p.A.	
		G	Procter & Gamble Pharmaceuticals France SAS.	
		G	Procter & Gamble Pharmaceuticals, Inc.	
		G	Procter & Gamble Pharmaceuticals Germany GmbH.	
		G	Procter & Gamble Pharmaceuticals S.a.r.l.	
		G	Procter & Gamble Pharmaceuticals Iberia S.L.	
20090752	G	G	Procter & Gamble Pharmaceuticals UK Limited.	
		G	LEO Fondet.	
		G	Peplin, Inc.	
		G	Peplin, Inc.	

FOR FURTHER INFORMATION CONTACT:

Sandra M. Peay, Contact Representative, or Renee Hallman, Contact Representative, Federal Trade Commission, Premerger Notification Office, Bureau of Competition, Room H-303, Washington, DC 20580, (202) 326-3100.

By direction of the Commission.

Donald S. Clark,

Secretary.

[FR Doc. E9-25376 Filed 10-21-09; 8:45 am]

BILLING CODE 6750-01-M

FEDERAL TRADE COMMISSION

Granting of Request for Early Termination of the Waiting Period Under the Premerger Notification Rules

Section 7A of the Clayton Act, 15 U.S.C. 18a, as added by Title II of the Hart-Scott Rodino Antitrust Improvements Act of 1976, requires persons contemplating certain mergers or acquisitions to give the Federal Trade Commission and the Assistant Attorney General advance notice and to wait designated periods before consummation of such plans. Section

7A(b)(2) of the Act permits the agencies, in individual cases, to terminate this waiting period prior to its expiration and requires that notice of this action be published in the **Federal Register**.

The following transactions were granted early termination of the waiting period provided by law and the premerger notification rules. The grants were made by the Federal Trade Commission and the Assistant Attorney General for the Antitrust Division of the Department of Justice. Neither agency intends to take any action with respect to these proposed acquisitions during the applicable waiting period.

TRANSACTION GRANTED EARLY TERMINATION

ET date	Trans No.	ET req status	Party name	
14-AUG-09	20090623	G	PartnerRe Ltd.	
		G	PARIS RE Holdings Limited.	
	20090627	G	PARIS RE Holdings Limited.	
		G	Oglethorpe Power Corporation.	
	20090630	G	G	International Power plc.
			G	Hartwell Energy Limited Partnership.
		G	G	Oglethorpe Power Corporation.
			G	Natural Gas Partners VIII, L.P.
	20090644	G	G	Hartwell Energy Limited Partnership.
			G	Career Education Corporation.

TRANSACTION GRANTED EARLY TERMINATION—Continued

ET date	Trans No.	ET req status	Party name	
18-AUG-09	20090648	G	Stitching LA Fondation Andre Cointreau.	
		G	Le Cordon Bleu International B.V.	
		G	McAfee, Inc.	
20-AUG-09	20090427	G	MX Logic, Inc.	
		G	MX Logic, Inc.	
		G	Arch Coal, Inc.	
21-AUG-09	20090448	G	Rio unto plc.	
		G	Jacobs Ranch Coal LLC.	
		G	Oracle Corporation.	
	20090655	G	Sun Microsystems, Inc.	
		G	Sun Microsystems, Inc.	
		G	Sprint Nextel Corporation.	
	20090657	G	Virgin Mobile USA, Inc.	
		G	Virgin Mobile USA, Inc.	
		G	ArcLight Energy Partners Fund III, L.P.	
	24-AUG-09	20090661	G	PPL Corporation.
			G	PPL Maine, LLC.
			G	ArcLight Energy Partners Fund IV, L.P.
20090665		G	PPL Corporation.	
		G	PPL Maine, LLC.	
		G	Targa Resources Partners LP.	
25-AUG-09		20090647	G	Targa Resources Investments Inc.
			G	Targa LSNG GP LLC.
			G	Targa LSNG LP.
	20090653	G	Targa Downstream GP LLC.	
		G	Targa Downstream LP.	
		G	Aetna Inc.	
28-AUG-09	20090654	G	Psychiatric Solutions, Inc.	
		G	Horizon Behavioral Services, LLC.	
		G	Manulife Financial Corporation.	
	20090664	G	PPL Corporation.	
		G	PPL Edgewood Energy, LLC.	
		G	PPL Shoreham Energy, LLC.	
	01-SEP-09	20090654	G	Electric Power Development Co., Ltd.
			G	PPL Corporation.
			G	PPL Edgewood Energy, LLC.
		20090664	G	PPL Shoreham Energy, LLC.
			G	Sentara Healthcare.
			G	Potomac Hospital Foundation.
02-SEP-09	20090645	G	Potomac Hospital Corporation of Prince William.	
		G	lochpe-Maxion S.A.	
		G	ArvinMeritor, Inc.	
	20090672	G	ArvinMeritor OE, LLC.	
		G	Meritor LVS S.A. de C.V.	
		G	Servicios Corporativos ArvinMeritor, S.A. de C.V.	
01-SEP-09	20090676	G	Meritor Comercio Industria de Sistemas Automotivos Ltda.	
		G	JPMorgan Chase & Co.	
		G	ArthroCare Corporation.	
	20090626	G	ArthroCare Corporation.	
		G	Noble Group Limited.	
		G	SemGroup, L.P.-Debtor-in-Possession.	
02-SEP-09	20090677	G	SemFuel, L.P.-Debtor-in-Possession.	
		G	Kurosawa B.V.	
		G	William B. Dunavant, Jr.	
	20090687	G	Dunavant Enterprises, Inc.	
		G	Frontier Communications Corporation.	
		G	Verizon Communications Inc.	
02-SEP-09	20090687	G	New Communications Holdings, Inc.	
		G	Inverness Medical Innovations, Inc.	
		G	Free & Clear, Inc.	
	20090679	G	Free & Clear, Inc.	
		G	LS Power Equity Partners II, L.P.	
		G	Dynegy, Inc.	
	20090680	G	Sandy Creek Services, LLC.	
		G	Riverside Generating Company, L.L.C.	
		G	Renaissance Power, LLC.	
G		Bridgeport Energy LLC.		
G		Bluegrass Generation Company, L.L.C.		
G		Dynegy Sandy Creek Holdings, LLC.		
	20090680	G	LS Power Equity Partners, L.P.	
		G	Dynegy, Inc.	

TRANSACTION GRANTED EARLY TERMINATION—Continued

ET date	Trans No.	ET req status	Party name
03-SEP-09	20090690	G	Tilton Energy LLC.
		G	Griffith Energy LLC.
		G	Dynegy Arlington Valley, LLC.
		G	Rocky Road Power, LLC.
		G	General Motors Company.
04-SEP-09	20090401	G	Delphi Corporation.
		G	DIP Holdco LLP.
		G	Fidelity National Information Services, Inc.
		G	Metavante Technologies, Inc.
		G	Metavante Technologies, Inc.
04-SEP-09	20090697	G	Electric Power Development Co., Ltd.
		G	General Electric Company.
		G	Birchwood Power Partners, L.P.
		G	Joe and Marlene Ricketts Grandchildren's Trust.
		G	Tribune Company.
04-SEP-09	20090702	G	Chicago Baseball Holdings, LLC.
		G	STG III, L.P.
		G	MSC Software Corporation.
		G	MSC Software Corporation.
		G	MSC Software Corporation.
04-SEP-09	20090704	G	MSC Software Corporation.
		G	MSC Software Corporation.
		G	MSC Software Corporation.
		G	MSC Software Corporation.
		G	MSC Software Corporation.

FOR FURTHER INFORMATION CONTACT:
Sandra M. Peay, Contact Representative,
or Renee Hallman, Contact
Representative, Federal Trade
Commission, Premerger Notification
Office, Bureau of Competition, Room H-
303, Washington, DC 20580, (202) 326-
3100.

By direction of the Commission.

Donald S. Clark,
Secretary.

[FR Doc. E9-25377 Filed 10-21-09; 8:45 am]

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**DEPARTMENT OF HEALTH AND
HUMAN SERVICES**

Food and Drug Administration

[Docket No. FDA-2009-N-0474]

**Agency Information Collection
Activities; Proposed Collection;
Comment Request; Inspection by
Accredited Persons Program Under
the Medical Device User Fee and
Modernization Act of 2002**

AGENCY: Food and Drug Administration,
HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing an
opportunity for public comment on the
proposed collection of certain
information by the agency. Under the
Paperwork Reduction Act of 1995 (the
PRA), Federal agencies are required to
publish notice in the **Federal Register**
concerning each proposed collection of
information, including each proposed
extension of an existing collection of
information, and to allow 60 days for

public comment in response to the
notice. This notice solicits comments on
the publication of the criteria FDA
intends to use to accredit third parties
to conduct inspections of eligible
manufacturers of class II or class III
medical devices.

DATES: Submit written or electronic
comments on the collection of
information by December 21, 2009.

ADDRESSES: Submit electronic
comments on the collection of
information to [http://
www.regulations.gov](http://www.regulations.gov). Submit written
comments on the collection of
information to the Division of Dockets
Management (HFA-305), Food and Drug
Administration, 5630 Fishers Lane, rm.
1061, Rockville, MD 20852. All
comments should be identified with the
docket number found in brackets in the
heading of this document.

FOR FURTHER INFORMATION CONTACT:
Denver Presley, Jr., Office of Information
Management (HFA-710), Food and Drug
Administration, 5600 Fishers Lane,
Rockville, MD 20857, 301-796-3793.

SUPPLEMENTARY INFORMATION: Under the
PRA (44 U.S.C. 3501-3520), Federal
agencies must obtain approval from the
Office of Management and Budget
(OMB) for each collection of
information they conduct or sponsor.
“Collection of information” is defined
in 44 U.S.C. 3502(3) and 5 CFR
1320.3(c) and includes agency requests
or requirements that members of the
public submit reports, keep records, or
provide information to a third party.
Section 3506(c)(2)(A) of the PRA (44
U.S.C. 3506(c)(2)(A)) requires Federal
agencies to provide a 60-day notice in
the **Federal Register** concerning each
proposed collection of information,

including each proposed extension of an
existing collection of information,
before submitting the collection to OMB
for approval. To comply with this
requirement, FDA is publishing notice
of the proposed collection of
information set forth in this document.

With respect to the following
collection of information, FDA invites
comments on these topics: (1) Whether
the proposed collection of information
is necessary for the proper performance
of FDA’s functions, including whether
the information will have practical
utility; (2) the accuracy of FDA’s
estimate of the burden of the proposed
collection of information, including the
validity of the methodology and
assumptions used; (3) ways to enhance
the quality, utility, and clarity of the
information to be collected; and (4)
ways to minimize the burden of the
collection of information on
respondents, including through the use
of automated collection techniques,
when appropriate, and other forms of
information technology.

**Inspection by Accredited Persons
Program Under the Medical Device
User Fee and Modernization Act of
2002; FD&C Act, Section 704(g) (OMB
Control Number 0910-0510)—Extension**

The Medical Device User Fee and
Modernization Act of 2002 (MDUFMA)
(Public Law 107-250) was signed into
law on October 26, 2002. Section 201 of
MDUFMA adds a new paragraph “g” to
section 704 of the Federal, Food, Drug,
and Cosmetic Act (the FD&C Act) (21
U.S.C. 374), directing FDA to accredit
third parties (accredited persons (APs))
to conduct inspections of eligible
manufacturers of class II or class III
devices. This is a voluntary program.