CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:

022063Orig1s000

ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS

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ACTION PACKAGE CHECKLIST

APPLICATION INFORMATION ¹						
NDA # 022063 BLA #	NDA Supplement # BLA Supplement #		If NDA, Efficacy Supplement Type: (an action package is not required for SE8 or SE9 supplements)			
Proprietary Name: Mydayis Established/Proper Name: mixed salts of a single-entity amphetamine product Dosage Form: Capsules			Applicant: Shire, LLC. Agent for Applicant (if applicable): Peggy Sung; psung@shire.com			
RPM: Latrice Wilson, PharmD; <u>latrice.wilson@fda.hhs.gov;</u> 240-402-5317			Division: Psychiatry Produ	cts		
NDA Application Type: 505(b)(1) 505(b)(2) Efficacy Supplement: 505(b)(1) 505(b)(2) BLA Application Type: 351(k) 351(a) Efficacy Supplement: 351(k) 351(a) Efficacy Supplement: 351(k) 351(a) Vote: Ij informa pediatri 0		Revie the d Chec exclu N N Date Note: If p informati	LL 505(b)(2) applications, two months prior to EVERY action: view the information in the 505(b)(2) Assessment and submit e draft ² to CDER OND IO for clearance. neck Orange Book for newly listed patents and/or clusivity (including pediatric exclusivity) No changes New patent/exclusivity (notify CDER OND IO) te of check: if pediatric exclusivity has been granted or the pediatric ation in the labeling of the listed drug changed, determine whether ic information needs to be added to or deleted from the labeling of tg.			
Actions						
 Proposed action User Fee Goal Date is June 20, 2017 			AP TA CR			
Previous actions (specify type and date for each action taken)			None Approvable 5/18/2007			
 If accelerated approval or approval based on efficacy studies in animals, were promotional materials received? Note: Promotional materials to be used within 120 days after approval must have been submitted (for exceptions, see http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/GuidanceSyncm069965.pdf). If not submitted, explain 			Received			
 Application Characteristics³ 						

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¹ The **Application Information** Section is (only) a checklist. The **Contents of Action Package** Section (beginning on page 2) lists the documents to be included in the Action Package.

 $^{^{2}}$ For resubmissions, 505(b)(2) applications must be cleared before the action, but it is not necessary to resubmit the draft 505(b)(2) Assessment to CDER OND IO unless the Assessment has been substantively revised (e.g., new listed drug, patent certification revised).

³ Answer all questions in all sections in relation to the pending application, i.e., if the pending application is an NDA or BLA supplement, then the questions should be answered in relation to that supplement, not in relation to the original NDA or BLA.

Fast Track Rx-to-OTC full switch Rolling Review Rx-to-OTC partial switch Orphan drug designation Direct-to-OTC Breakthrough Therapy designation Direct-to-OTC (NOTE: Set the submission property in DARRTS and notify the CDER Breakthrough Therapy Program Manager;				
(<u>NOTE</u> , Set the submission property in DARKI'S and holy the CDER Breakinrough Therapy Program Manager, Refer to the "RPM BT Checklist for Considerations after Designation Granted" for other required actions: <u>CST SharePoint</u>)				
NDAs: Subpart H BLAs: Subpart E Accelerated approval (21 CFR 314.510) Accelerated approval (21 CFR 314.520) Subpart I Reproval based on animal studies Approval based on animal studies Approval based on animal studies				
 Submitted in response to a PMR Submitted in response to a PMC Submitted in response to a Pediatric Written Request Submitted in response to a Pediatric Written Request REMS: MedGuide MedGuide w/o REMS REMS not required 				
Comments:				
 ◆ BLAs only: Is the product subject to official FDA lot release per 21 CFR 610.2 (approvals only) □ Yes □ No 				
Public communications (approvals only)				
Office of Executive Programs (OEP) liaison has been notified of action Yes No				
 Indicate what types (if any) of information were issued Indicate what types (if any) of information were issued FDA Talk Paper CDER Q&As Other 				
Exclusivity				
 Is approval of this application blocked by any type of exclusivity (orphan, 5-year NCE, 3-year, pediatric exclusivity)? If so, specify the type 				
✤ Patent Information (NDAs only)				
 Patent Information: Verify that form FDA-3542a was submitted for patents that claim the drug for which approval is sought. Not applicable because drug an old antibiotic. 	is			
CONTENTS OF ACTION PACKAGE				
Officer/Employee List (Tab A)				
 List of officers/employees who participated in the decision to approve this application and consented to be identified on this list (approvals only) Included 				
Documentation of consent/non-consent by officers/employees				

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Action Letters (Tab B)						
*	Copies of all action letters (including approval letter with final labeling)	Action(s) and date(s): Approval 6/20/2017 Approvable 05/18/2007				
	Labeling (Tab C)					
*	Package Insert (write submission/communication date at upper right of first page of PI)					
	 Most recent draft labeling (if it is division-proposed labeling, it should be in track-changes format) 	Included				
	Original applicant-proposed labeling	Included				
*	Medication Guide/Patient Package Insert/Instructions for Use/Device Labeling (write submission/communication date at upper right of first page of each piece)	 Medication Guide Patient Package Insert Instructions for Use Device Labeling None 				
	 Most-recent draft labeling (if it is division-proposed labeling, it should be in track-changes format) 	Included				
	Original applicant-proposed labeling	Included				
*	Labels (full color carton and immediate-container labels) (write submission/communication date on upper right of first page of each submission)					
	Most-recent draft labeling	Included				
*	 Proprietary Name Acceptability/non-acceptability letter(s) (indicate date(s)) Review(s) (indicate date(s) 	04/14/2017 04/13/2017				
*	Labeling reviews (indicate dates of reviews)	RPM: None DMEPA: None 5/18/2017; 6/19/2017; 6/20/2017 0/19/2017; DMPP/PLT (DRISK): □ □ None 5/30/2017 OPDP: □ None 05/26/2017; 5/30/2017 SEALD: ○ None CSS: ○ None Orduct Quality ○ None Other:				
Administrative / Regulatory Documents (Tab D)						
* *	RPM Filing Review ⁴ /Memo of Filing Meeting <i>(indicate date of each review)</i> All NDA 505(b)(2) Actions: Date each action cleared by 505(b)(2) Clearance Committee	N/A Not a (b)(2)				
*	NDAs/NDA supplements only: Exclusivity Summary (signed by Division Director)	Completed (Do not include)				
*	Application Integrity Policy (AIP) Status and Related Documents http://www.fda.gov/ICECI/EnforcementActions/ApplicationIntegrityPolicy/default.htm					
	Applicant is on the AIP	🗌 Yes 🛛 No				

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	This application is on the AIP	🗌 Yes 🕅 No		
	• If yes, Center Director's Exception for Review memo (indicate date)			
	• If yes, OC clearance for approval (indicate date of clearance communication)	☐ Not an AP action		
*	 Pediatrics (approvals only) Date reviewed by PeRC <u>5/17/2017</u> If PeRC review not necessary, explain: 			
*	Breakthrough Therapy Designation	N/A		
	Breakthrough Therapy Designation Letter(s) (granted, denied, an/or rescinded)			
	 CDER Medical Policy Council Breakthrough Therapy Designation Determination Review Template(s) (include only the completed template(s) and not the meeting minutes) 			
	 CDER Medical Policy Council Brief – Evaluating a Breakthrough Therapy Designation for Rescission Template(s) (include only the completed template(s) and not the meeting minutes) (completed CDER MPC templates can be found in DARRTS as clinical reviews or on the MPC SharePoint Site) 			
*	Outgoing communications: letters, emails, and faxes considered important to include in the action package by the reviewing office/division (e.g., clinical SPA letters, RTF letter, Formal Dispute Resolution Request decisional letters, etc.) (do not include OPDP letters regarding pre-launch promotional materials as these are non-disclosable; do not include Master File letters; do not include previous action letters, as these are located elsewhere in package)	Ack Letter: 1/17/2017 Information Requests: 1/17/2017; 3/13/2017; 5/25/2017; PMR: 5/15/2017 Approval Receipt: 6/20/2017		
*	Internal documents: memoranda, telecons, emails, and other documents considered important to include in the action package by the reviewing office/division (e.g., Regulatory Briefing minutes, Medical Policy Council meeting minutes)			
*	Minutes of Meetings			
	• If not the first review cycle, any end-of-review meeting (indicate date of mtg)	⊠ N/A or no mtg		
	Pre-NDA/BLA meeting (indicate date of mtg)	🛛 No mtg		
	• EOP2 meeting (indicate date of mtg)	No mtg		
	Mid-cycle Communication (indicate date of mtg)	N/A		
	Late-cycle Meeting (indicate date of mtg)	N/A		
	 Other milestone meetings (e.g., EOP2a, CMC focused milestone meetings) (indicate dates of mtgs) 	N/A		
*	Advisory Committee Meeting(s)	No AC meeting		
	• Date(s) of Meeting(s)			
Decisional and Summary Memos (Tab E)				
*	Office Director Decisional Memo (indicate date for each review)	🛛 None		
	Division Director Summary Review (indicate date for each review)	None 6/19/2017		
	Cross-Discipline Team Leader Review (indicate date for each review)	None 6/19/2017		
	PMR/PMC Development Templates (indicate total number)	None 4		
Clinical (Tab F)				

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