CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:

213736Orig1s000

PROPRIETARY NAME REVIEW(S)

PROPRIETARY NAME REVIEW

Division of Medication Error Prevention and Analysis (DMEPA)
Office of Medication Error Prevention and Risk Management (OMEPRM)
Office of Surveillance and Epidemiology (OSE)
Center for Drug Evaluation and Research (CDER)

*** This document contains proprietary information that cannot be released to the public***

Date of This Review: December 13, 2019

Application Type and Number: NDA 213736

Product Name and Strength: Pemazyre (pemigatinib) Tablets,

4.5 mg, 9 mg and 13.5 mg

Product Type: Single Ingredient Product

Rx or OTC: Prescription (Rx)

Applicant/Sponsor Name: Incyte Corporation (Incyte)

Panorama #: 2019-34840153

DMEPA Safety Evaluator: Maximilian Straka, PharmD, FISMP

DMEPA Team Leader: Chi-Ming (Alice) Tu, PharmD, FISMP, BCPS

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1 INTRODUCTION

This review evaluates the proposed proprietary name, Pemazyre, from a safety and misbranding perspective. The sources and methods used to evaluate the proposed proprietary name are outlined in the reference section and Appendix A respectively. Incyte submitted an external name study, conducted for this proposed proprietary name. The external study was previously evaluated by DMEPA (See section 1.1).^a

1.1 REGULATORY HISTORY

Incyte previously submitted the proposed proprietary name, Pemazyre*** on April 5, 2018. We found the name, Pemazyre, conditionally acceptable on September 26, 2018 under IND 138179.^a

Subsequently, Incyte submitted the proposed proprietary name, Pemazyre under NDA 213736, for review on October 1, 2019. We note that the product characteristics submitted on October 1, 2019 have changed (dose reduction information has been added) since our last IND review.

1.2 PRODUCT INFORMATION

The following product information is provided in the proprietary name submission received on October 1, 2019.

- Intended Pronunciation: pem' ah zeer
- Active Ingredient: pemigatinib
- Indication of Use: The treatment of adults with previously treated, locally advanced or metastatic cholangiocarcinoma with a fibroblast growth factor receptor 2 (FGFR2) fusion or rearrangement.
- Route of Administration: Oral
- Dosage Form: Tablets
- Strength: 4.5 mg, 9 mg and 13.5 mg
- Dose and Frequency:
 - The recommended dose of PEMAZYRE is 13.5 mg taken orally once daily for 14 days followed by 7 days off therapy. Continue treatment until disease progression or unacceptable toxicity occurs.
 - o The recommended dose reductions are:
 - First dose reduction: PEMAZYRE 9 mg taken orally once daily for 14 days on, followed by 7 days off therapy.
 - Second dose reduction: PEMAZYRE 4.5 mg taken orally once daily for 14 days on, followed by 7 days off therapy.

^a Little, C. Proprietary Name Review for Pemazyre (IND 138179). Silver Spring (MD): FDA, CDER, OSE, DMEPA (US); 2018 SEP 26. Panorama No. 2018-22138081.

- Permanently discontinue if unable to tolerate PEMAZYRE 4.5 mg once daily.
- How Supplied: PEMAZYRE (pemigatinib) tablets are available as follows:
 - o 4.5 mg tablets: Round, white to off-white debossed on one side with "I" and "4.5" on the other side in bottles of 14 with child-resistant closure, NDC 50881-026-01
 - o 9 mg tablets: Oval, white to off-white debossed on one side with "I" and "9" on the other side in bottles of 14 with child-resistant closure, NDC 50881-027-01
 - 13.5 mg tablets: Round, white to off-white debossed on one side with "I" and
 "13.5" on the other side in bottles of 14 with child-resistant closure, NDC 50881-028-01
- Storage: Store PEMAZYRE bottles at room temperature 20 25°C (68 77°F); excursions permitted to 15 30°C (59 86°F).

2 RESULTS

The following sections provide information obtained and considered in the overall evaluation of the proposed proprietary name, Pemazyre.

2.1 MISBRANDING ASSESSMENT

The Office of Prescription Drug Promotion (OPDP) determined that Pemazyre would not misbrand the proposed product. The Division of Medication Error Prevention and Analysis (DMEPA) and the Division of Oncology Products 2 (DOP2) concurred with the findings of OPDP's assessment for Pemazyre.

2.2 SAFETY ASSESSMENT

The following aspects were considered in the safety evaluation of the proposed proprietary name, Pemazyre.

2.2.1 United States Adopted Names (USAN) Search

There is no USAN stem present in the proposed proprietary name^b.

2.2.2 Components of the Proposed Proprietary Name

Incyte indicated in their submission that the proposed name, Pemazyre, contains a prefix (PEM) linked to the established name pemigatinib with a blank canvas suffix. This proprietary name is comprised of a single word that does not contain any components (i.e. a modifier, route of administration, dosage form, etc.) that are misleading or can contribute to medication error.

2.2.3 Comments from Other Review Disciplines at Initial Review

In response to the OSE, October 21, 2019 e-mail, the Division of Oncology Products 2 (DOP2) did not forward any comments or concerns relating to Pemazyre at the initial phase of the review.

^b USAN stem search conducted on October 24, 2019.

2.2.4 FDA Name Simulation Studies

Seventy-three (73) practitioners participated in DMEPA's prescription studies for Pemazyre. The responses did not overlap with any currently marketed products nor did the responses sound or look similar to any currently marketed products or any products in the pipeline. Appendix B contains the results from the verbal and written prescription studies.

2.2.5 Phonetic and Orthographic Computer Analysis (POCA) Search Results

Our POCA search^c identified 103 names with the combined score of ≥55% or individual orthographic or phonetic score of ≥70%. We had identified and evaluated some of the names in our previous proprietary name review.^a We re-evaluated the previously identified names of concern considering any lessons learned from recent post-marketing experience, which may have altered our previous conclusion regarding the acceptability of the name. We note that in our previous proprietary name review, we evaluated the proposed dose of "13.5 mg orally once daily for two weeks, followed by one week off therapy." In this current submission, Incyte included two dose reductions: 1. "PEMAZYRE 9 mg taken orally once daily for 14 days on, followed by 7 days off therapy" and 2. "PEMAZYRE 4.5 mg taken orally once daily for 14 days on, followed by 7 days off therapy." We find that these changes to the product characteristics do not alter our assessment of the names evaluated in our previous review, and we agree with our previous findings. Therefore, we identified 5 names not previously analyzed. These names are included in Table 1 below.

2.2.6 Names Retrieved for Review Organized by Name Pair Similarity

Table 1 lists the number of names retrieved from our POCA search. These name pairs are organized as highly similar, moderately similar or low similarity for further evaluation.

Table 1. Names Retrieved for Review Organized by Name Pair Similarity			
Similarity Category	Number of Names		
Highly similar name pair: combined match percentage score ≥70%	1		
Moderately similar name pair: combined match percentage score ≥55% to ≤ 69%	4		
Low similarity name pair: combined match percentage score ≤54%	0		

2.2.7 Safety Analysis of Names with Potential Orthographic, Spelling, and Phonetic Similarities

Our analysis of the 5 names contained in Table 1 determined none of the names will pose a risk for confusion with Pemazyre as described in Appendices C through H.

^c POCA search conducted on October 24, 2019 in version 4.3.

2.2.8 Communication of DMEPA's Analysis at Midpoint of Review

DMEPA communicated our findings to the Division of Oncology 3 (DO3) via e-mail on November 25, 2019. At that time we also requested additional information or concerns that could inform our review. Per e-mail correspondence from DO3 on December 2, 2019, they stated the proposed proprietary name Pemazyre sounds like pemetrexed in general and Pemfexy***. Both "pemetrexed" and "Pemfexy***" were included for evaluation in our previous proprietary name review, where we concluded both pemetrexed and Pemfexy*** do not sound-like the proposed proprietary name Pemazyre.^a Taking into account the comments from DO3, we reevaluated the names and we maintain our previous findings.

3 CONCLUSION

The proposed proprietary name, Pemazyre, is acceptable.

If you have any questions or need clarifications, please contact Latonia Ford, OSE project manager, at 301-796-4901.

3.1 COMMENTS TO INCYTE CORPORATION

We have completed our review of the proposed proprietary name, Pemazyre, and have concluded that this name is acceptable.

If any of the proposed product characteristics as stated in your submission, received on October 1, 2019, are altered prior to approval of the marketing application, the name must be resubmitted for review.

REFERENCES

USAN Stems (<u>https://www.ama-assn.org/about/united-states-adopted-names-approved-stems</u>)
 USAN Stems List contains all the recognized USAN stems.

2. Phonetic and Orthographic Computer Analysis (POCA)

POCA is a system that FDA designed. As part of the name similarity assessment, POCA is used to evaluate proposed names via a phonetic and orthographic algorithm. The proposed proprietary name is converted into its phonemic representation before it runs through the phonetic algorithm. Likewise, an orthographic algorithm exists that operates in a similar fashion. POCA is publicly accessible.

Drugs@FDA

Drugs@FDA is an FDA Web site that contains most of the drug products approved in the United States since 1939. The majority of labels, approval letters, reviews, and other information are available for drug products approved from 1998 to the present. Drugs@FDA contains official information about FDA-approved *brand name* and *generic drugs*; *therapeutic biological products*, *prescription* and *over-the-counter* human drugs; and *discontinued drugs* (see Drugs @ FDA Glossary of Terms, available at http://www.fda.gov/Drugs/InformationOnDrugs/ucm079436.htm#ther-biological).

RxNorm

RxNorm contains the names of prescription and many OTC drugs available in the United States. RxNorm includes generic and branded:

- Clinical drugs pharmaceutical products given to (or taken by) a patient with therapeutic or diagnostic intent
- Drug packs packs that contain multiple drugs, or drugs designed to be administered in a specified sequence

Radiopharmaceuticals, contrast media, food, dietary supplements, and medical devices, such as bandages and crutches, are all out of scope for RxNorm (http://www.nlm.nih.gov/research/umls/rxnorm/overview.html).

Division of Medication Errors Prevention and Analysis proprietary name consultation requests

This is a list of proposed and pending names that is generated by the Division of Medication Error Prevention and Analysis from the Access database/tracking system.

APPENDICES

Appendix A

FDA's Proprietary Name Risk Assessment evaluates proposed proprietary names for misbranding and safety concerns.

- 1. **Misbranding Assessment**: For prescription drug products, OPDP assesses the name for misbranding concerns. For over-the-counter (OTC) drug products, the misbranding assessment of the proposed name is conducted by DNDP. OPDP or DNDP evaluates proposed proprietary names to determine if the name is false or misleading, such as by making misrepresentations with respect to safety or efficacy. For example, a fanciful proprietary name may misbrand a product by suggesting that it has some unique effectiveness or composition when it does not (21 CFR 201.10(c)(3)). OPDP or DNDP provides their opinion to DMEPA for consideration in the overall acceptability of the proposed proprietary name.
- 2. **Safety Assessment**: The safety assessment is conducted by DMEPA, and includes the following:
- a. Preliminary Assessment: We consider inclusion of USAN stems or other characteristics that when incorporated into a proprietary name may cause or contribute to medication errors (i.e., dosing interval, dosage form/route of administration, medical or product name abbreviations, names that include or suggest the composition of the drug product, etc.) See prescreening checklist below in Table 2*. DMEPA defines a medication error as any preventable event that may cause or lead to inappropriate medication use or patient harm while the medication is in the control of the health care professional, patient, or consumer. d

^d National Coordinating Council for Medication Error Reporting and Prevention. http://www.nccmerp.org/aboutMedErrors.html. Last accessed 10/11/2007.

*Table 2- Prescreening Checklist for Proposed Proprietary Name

	Answer the questions in the checklist below. Affirmative answers to any of these questions indicate a potential area of concern that should be carefully evaluated as described in this guidance.				
Y/N	Is the proposed name obviously similar in spelling and pronunciation to other names?				
	Proprietary names should not be similar in spelling or pronunciation to proprietary names, established names, or ingredients of other products.				
Y/N	Are there inert or inactive ingredients referenced in the proprietary name?				
	Proprietary names should not incorporate any reference to an inert or inactive ingredient in a way that might create an impression that the ingredient's value is greater than its true functional role in the formulation (21 CFR 201.10(c)(4)).				
Y/N	Does the proprietary name include combinations of active ingredients?				
	Proprietary names of fixed combination drug products should not include or suggest the name of one or more, but not all, of its active ingredients (see 21 CFR 201.6(b)).				
Y/N	Is there a United States Adopted Name (USAN) stem in the proprietary name?				
	Proprietary names should not incorporate a USAN stem in the position that USAN designates for the stem.				
Y/N	Is this proprietary name used for another product that does not share at least one common active ingredient?				
	Drug products that do not contain at least one common active ingredient should not use the same (root) proprietary name.				
Y/N	Is this a proprietary name of a discontinued product?				
	Proprietary names should not use the proprietary name of a discontinued product if that discontinued drug product does not contain the same active ingredients.				

- b. Phonetic and Orthographic Computer Analysis (POCA): Following the preliminary screening of the proposed proprietary name, DMEPA staff evaluates the proposed name against potentially similar names. In order to identify names with potential similarity to the proposed proprietary name, DMEPA enters the proposed proprietary name in POCA and queries the name against the following drug reference databases, Drugs@fda, CernerRxNorm, and names in the review pipeline using a 55% threshold in POCA. DMEPA reviews the combined orthographic and phonetic matches and group the names into one of the following three categories:
 - Highly similar pair: combined match percentage score $\geq 70\%$.
 - Moderately similar pair: combined match percentage score \geq 55% to \leq 69%.

• Low similarity: combined match percentage score ≤54%.

Using the criteria outlined in the check list (Table 3-5) that corresponds to each of the three categories (highly similar pair, moderately similar pair, and low similarity), DMEPA evaluates the name pairs to determine the acceptability or non-acceptability of a proposed proprietary name. The intent of these checklists is to increase the transparency and predictability of the safety determination of whether a proposed name is vulnerable to confusion from a look-alike or sound-alike perspective. Each bullet below corresponds to the name similarity category cross-references the respective table that addresses criteria that DMEPA uses to determine whether a name presents a safety concern from a look-alike or sound-alike perspective.

- For highly similar names, differences in product characteristics often cannot mitigate the risk of a medication error, including product differences such as strength and dose. Thus, proposed proprietary names that have a combined score of ≥ 70 percent are at risk for a look-alike sound-alike confusion which is an area of concern (See Table 3).
- Moderately similar names are further evaluated to identify the presence of attributes that are known to cause name confusion.
 - Name attributes: We note that the beginning of the drug name plays a significant role in contributing to confusion. Additionally, drug name pairs that start with the same first letter and contain a shared letter string of at least 3 letters in both names are major contributing factor in the confusion of drug names. We evaluate all moderately similar names retrieved from POCA to identify the above attributes. These names are further evaluated to identify overlapping or similar strengths or doses.
 - Product attributes: Moderately similar names of products that have overlapping or similar strengths or doses represent an area for concern for FDA. The dose and strength information is often located in close proximity to the drug name itself on prescriptions and medication orders, and the information can be an important factor that either increases or decreases the potential for confusion between similarly named drug pairs. The ability of other product characteristics to mitigate confusion (e.g., route, frequency, dosage form) may be limited when the strength or dose overlaps. DMEPA reviews such names further, to determine whether sufficient differences exist to prevent confusion. (See Table 4).
- Names with low similarity that have no overlap or similarity in strength and dose are generally acceptable (See Table 5) unless there are data to suggest that the name might be vulnerable to confusion (e.g., prescription simulation study suggests that the name is likely to be misinterpreted as a marketed product). In these instances, we would reassign

^e Shah, M, Merchant, L, Characteristics That May Help in the Identification of Potentially Confusing Proprietary Drug Names. Therapeutic Innovation & Regulatory Science, September 2016

a low similarity name to the moderate similarity category and review according to the moderately similar name pair checklist.

c. FDA Prescription Simulation Studies: DMEPA staff also conducts a prescription simulation studies using FDA health care professionals.

Three separate studies are conducted within the Centers of the FDA for the proposed proprietary name to determine the degree of confusion of the proposed proprietary name with marketed U.S. drug names (proprietary and established) due to similarity in visual appearance with handwritten prescriptions or verbal pronunciation of the drug name. The studies employ healthcare professionals (pharmacists, physicians, and nurses), and attempts to simulate the prescription ordering process. The primary Safety Evaluator uses the results to identify orthographic or phonetic vulnerability of the proposed name to be misinterpreted by healthcare practitioners.

In order to evaluate the potential for misinterpretation of the proposed proprietary name in handwriting and verbal communication of the name, inpatient medication orders and/or outpatient prescriptions are written, each consisting of a combination of marketed and unapproved drug products, including the proposed name. These orders are optically scanned and one prescription is delivered to a random sample of participating health professionals via e-mail. In addition, a verbal prescription is recorded on voice mail. The voice mail messages are then sent to a random sample of the participating health professionals for their interpretations and review. After receiving either the written or verbal prescription orders, the participants record their interpretations of the orders which are recorded electronically.

d. Comments from Other Review Disciplines: DMEPA requests the Office of New Drugs (OND) and/or Office of Generic Drugs (OGD), ONDQA or OBP for their comments or concerns with the proposed proprietary name, ask for any clinical issues that may impact the DMEPA review during the initial phase of the name review. Additionally, when applicable, at the same time DMEPA requests concurrence/non-concurrence with OPDP's decision on the name. The primary Safety Evaluator addresses any comments or concerns in the safety evaluator's assessment.

The OND/OGD Regulatory Division is contacted a second time following our analysis of the proposed proprietary name. At this point, DMEPA conveys their decision to accept or reject the name. The OND or OGD Regulatory Division is requested to provide any further information that might inform DMEPA's final decision on the proposed name.

Additionally, other review disciplines opinions such as ONDQA or OBP may be considered depending on the proposed proprietary name.

When provided, DMEPA considers external proprietary name studies conducted by or for the Applicant/Sponsor and incorporates the findings of these studies into the overall risk assessment. The DMEPA primary reviewer assigned to evaluate the proposed proprietary name is responsible for considering the collective findings, and provides an overall risk assessment of the proposed proprietary name.

Table 3. Highly Similar Name Pair Checklist (i.e., combined Orthographic and Phonetic score is $\geq 70\%$).

Answer the questions in the checklist below. Affirmative answers to some of these questions suggest that the pattern of orthographic or phonetic differences in the names may render the names less likely to confusion, provided that the pair does not share a common strength or dose.

Orthographic Checklist Phonetic Checklist			
	Statograpine Checkinst	I nonette encentist	
Y/N	Do the names begin with different first letters?	Y/N	Do the names have different number of syllables?
	Note that even when names begin with different first letters, certain letters may be confused with each other when scripted.		
Y/N	Are the lengths of the names dissimilar* when scripted?	Y/N	Do the names have different syllabic stresses?
	*FDA considers the length of names different if the names differ by two or more letters.		
Y/N	Considering variations in scripting of some letters (such as <i>z</i> and <i>f</i>), is there a different number or placement of upstroke/downstroke letters present in the names?	Y/N	Do the syllables have different phonologic processes, such vowel reduction, assimilation, or deletion?
Y/N	Is there different number or placement of cross-stroke or dotted letters present in the names?	Y/N	Across a range of dialects, are the names consistently pronounced differently?
Y/N	Do the infixes of the name appear dissimilar when scripted?		
Y/N	Do the suffixes of the names appear dissimilar when scripted?		

Table 4: Moderately Similar Name Pair Checklist (i.e., combined score is $\geq 55\%$ to $\leq 69\%$).

Step 1 Review the DOSAGE AND ADMINISTRATION and HOW SUPPLIED/STORAGE AND HANDLING sections of the prescribing information (or for OTC drugs refer to the Drug Facts label) to determine if strengths and doses of the name pair overlap or are very similar. Different strengths and doses for products whose names are moderately similar may decrease the risk of confusion between the moderately similar name pairs. Name pairs that have overlapping or similar strengths or doses have a higher potential for confusion and should be evaluated further (see Step 2). Because the strength or dose could be used to express an order or prescription for a particular drug product, overlap in one or both of these components would be reason for further evaluation.

For single strength products, also consider circumstances where the strength may not be expressed.

For any i.e. drug products comprised of more than one active ingredient, consider whether the strength or dose may be expressed using only one of the components.

To determine whether the strengths or doses are similar to your proposed product, consider the following list of factors that may increase confusion:

- Alternative expressions of dose: 5 mL may be listed in the prescribing information, but the dose may be expressed in metric weight (e.g., 500 mg) or in non-metric units (e.g., 1 tsp, 1 tablet/capsule). Similarly, a strength or dose of 1000 mg may be expressed, in practice, as 1 g, or vice versa.
- Trailing or deleting zeros: 10 mg is similar in appearance to 100 mg which may potentiate confusion between a name pair with moderate similarity.
- Similar sounding doses: 15 mg is similar in sound to 50 mg

Step 2 Answer the questions in the checklist below. Affirmative answers to some of these questions suggest that the pattern of orthographic or phonetic differences in the names may reduce the likelihood of confusion for moderately similar names with overlapping or similar strengths or doses.

Orthographic Checklist (Y/N to each question)

- Do the names begin with different first letters?
 - Note that even when names begin with different first letters, certain letters may be confused with each other when scripted.
- Are the lengths of the names dissimilar* when scripted?
 *EDA considers the length of names
 - *FDA considers the length of names different if the names differ by two or more letters.
- Considering variations in scripting of some letters (such as *z* and *f*), is there a different number or placement of upstroke/downstroke letters present in the names?
- Is there different number or placement of cross-stroke or dotted letters present in the names?
- Do the infixes of the name appear dissimilar when scripted?
- Do the suffixes of the names appear dissimilar when scripted?

Phonetic Checklist (Y/N to each question)

- Do the names have different number of syllables?
- Do the names have different syllabic stresses?
- Do the syllables have different phonologic processes, such vowel reduction, assimilation, or deletion?
- Across a range of dialects, are the names consistently pronounced differently?

Table 5: Low Similarity Name Pair Checklist (i.e., combined score is ≤54%).

Names with low similarity are generally acceptable unless there are data to suggest that the name might be vulnerable to confusion (e.g., prescription simulation study suggests that the name is likely to be misinterpreted as a marketed product). In these instances, we would reassign a low similarity name to the moderate similarity category and review according to the moderately similar name pair checklist.

Appendix B: Prescription Simulation Samples and Results

Figure 1. Pemazyre Study (Conducted on October 18, 2019)

Handwritten Medication Order/Prescription	Verbal Prescription
Medication Order:	Pemazyre 13.5 mg
Pemerzyne 9 mg po QD	Take 1 tablet by mouth daily for 14 days
Outpatient Prescription:	Dispense # 14
Pemazyre 13.5 mg — tabpo daily x 14 days # 14	

FDA Prescription Simulation Responses (Aggregate Report)

OUTPATIENT

As of Date 11/1/2019

215 People Received Study73 People Responded

TOTAL

INPATIENT

VOICE

Study Name: Pemazyre

Total

INTERPRETATION

TEMAZERE

TEMAZIR

TEMAZYR

TEMEVIR

TEMEZERE

TEMIZIER

TEMOVIR

PEMAZYME	0	0	1	1
PEMAZYN	0	0	1	1
PEMAZYNE	2	0	1	3
PEMAZYRE	35	0	13	48
PEMAZYRE 13.5 MG	1	0	0	1
PEMAZYU	0	0	1	1
PEMERZEPRE	0	0	1	1
PENAZYNE	1	0	0	1
PENAZYRE	2	0	1	3
TAMNISYR	0	1	0	1
TEMAZEER	0	2	0	2

Appendix C: Highly Similar Names (e.g., combined POCA score is $\geq 70\%$)

No.	Proposed name: Pemazyre	POCA	Orthographic and/or phonetic
	Established name: pemigatinib	Score (%)	differences in the names sufficient to
	Dosage form: Tablets		prevent confusion
	Strength(s): 4.5 mg, 9 mg and		
	13.5 mg		Other prevention of failure mode
	Usual Dose: One tablet by		expected to minimize the risk of
	mouth once daily for 14 days		confusion between these two names.
	followed by 7 days off therapy.		
1.	Pemazyre***	100	This name is the subject of this review.

Appendix D: Moderately Similar Names (e.g., combined POCA score is \geq 55% to \leq 69%) with no overlap or numerical similarity in Strength and/or Dose

No.	Name	POCA Score (%)
	N/A	

Appendix E: Moderately Similar Names (e.g., combined POCA score is $\geq 55\%$ to $\leq 69\%$) with overlap or numerical similarity in Strength and/or Dose

No.	Proposed name: Pemazyre	POCA	Prevention of Failure Mode
	Established name: pemigatinib	Score (%)	
	Dosage form: Tablets		In the conditions outlined below, the
	Strength(s): 4.5 mg, 9 mg and		following combination of factors, are
	13.5 mg		expected to minimize the risk of
	Usual Dose: One tablet by		confusion between these two names
	mouth once daily for 14 days		
	followed by 7 days off therapy.		
2.	Panzyga	58	This name pair has sufficient
			orthographic and phonetic differences.
3.	Zepsyre***	64	This name pair has sufficient
	T and the second	l .	orthographic and phonetic differences.

Appendix F: Low Similarity Names (e.g., combined POCA score is ≤54%)

No.	Name	POCA Score (%)
	N/A	

Appendix G: Names not likely to be confused or not used in usual practice settings for the reasons described.

No.	Name	POCA Score (%)	Failure preventions
	N/A		

Appendix H: Names not likely to be confused due to absence of attributes that are known to cause name confusion^f.

No.	Name	POCA Score (%)
4.	Dermacerin	56
5.	Zemdri	55

 $^{^{\}rm f}$ Shah, M, Merchant, L, Chan, I, and Taylor, K. Characteristics That May Help in the Identification of Potentially Confusing Proprietary Drug Names. Therapeutic Innovation & Regulatory Science, September 2016

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