

IN THE UNITED STATES DISTRICT COURT
FOR THE NORTHERN DISTRICT OF CALIFORNIA
SAN JOSE DIVISION

Margarita A.G., a minor Gaeta, child, as guardian et al.,
ad litem for v. Plaintiffs, Perrigo Pharmaceuticals Company, et al.,

Defendants. /

NO. C 05-04115 JW

ORDER PERRIGO’S JUDGMENT GRANTING MOTION DEFENDANT FOR

SUMMARY I. INTRODUCTION

Margarita Gaeta and Augustine Gaeta (collectively, “Plaintiffs”), bring this diversity action on behalf of their son, A.G., against Perrigo Pharmaceuticals Company (“Perrigo”), PAR Pharmaceutical Inc. (“PAR”), and BASF Corporation (“BASF”) (collectively, “Defendants”), alleging, *inter alia*, strict products liability, breach of warranty, and negligence. Plaintiffs allege A.G. suffered liver failure as a result of his consumption of ibuprofen manufactured and distributed

by Defendants. Presently before the Court is Defendant Perrigo’s Motion for Summary

Judgment.

(hereafter, “Motion,” Docket Item No. 156.) The Court conducted a hearing on April 14, 2008.

Based on the papers submitted to date and oral arguments of counsel, the Court GRANTS

Defendant

Perrigo’s Motion for Summary Judgment.

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igo’s generic over-the-counter (“OTC”) ibuprofen at
to June 6, 2004, A.G. took 400mg of the ibuprofen
, A.G. developed a fever, and he was seen by his
prescription-strength ibuprofen (400mg) to him.
ever, A.G.’s condition continued to worsen: on June 13,
m with a diagnosis of septic shock, dehydration,
transferred to Stanford University Hospital for a liver
04. (Id., Ex. A at 4-6.) A.G. developed other
ve necrotic tissue on his fingers and toes amputated. (Id.,

Role in Regulating Drugs

ics Act (“FDCA”), 21 U.S.C. §§ 301 *et seq.*, a

II. BACKGROUND

administration (“FDA”) approval before a new drug

and two benign moles removed in a surgical procedure. (Second U.S.C. § 355. The process for approval requires

after, “SAC,” Docket Item No. 29.) During the procedure, A.G.’s

Halothane, an anesthetic known to cause liver failure in certain

Z & LAURIE D. DELEVE, DRUG-INDUCED LIVER DISEASE 406-07 for Summary Judgment, Ex. A at 3, hereafter,

er the surgery, A.G. was discharged with instruction to take 400mg ^{iffs} have moved for leave to file a supplemental reply.

rs as needed for pain. (Declaration of Kelly J. Savage, hereafter, proffered supplemental reply is entirely duplicative of

-117:7, Docket Item No.

1 submission of a new drug application (“NDA”) to demonstrate that the drug is “safe and effective.”
2 21 U.S.C. § 355(a)-(i). The proof of the efficacy and safety of the drug must be based on extensive
3 laboratory testing. 21 U.S.C. § 355(b). Drug manufacturers also must submit to the FDA
4 “specimens of the labeling proposed to be used for such drug.” 21 U.S.C. § 355(b)(1). The label
5 must contain a “warnings” section which describes “clinically significant adverse reactions
6 (including any that are potentially fatal, are serious even if infrequent, or can be prevented or
7 mitigated through appropriate use of the drug).” 21 C.F.R. § 201.57(c)(6)(I). When the drug is
8 approved, the FDA includes it in its published list of approved drugs. See 21 U.S.C. § 355(j)(7).
9 The drug is then referred to as a “listed drug.” Id. § 355(j)(2)(A)(i). A listed drug may also be
10 referred to as an “innovator” or “pioneer” drug. See, e.g., *Bristol-Myers Squibb Co. v. Shalala*, 91
11 F.3d 1493, 1494, 1497-98 (D.C. Cir. 1996).
12 Before 1984, generic drug manufacturers were required to submit their own NDA. See
13 *Tri-Bio Labs., Inc. v. United States*, 836 F.2d 135, 138-39 (3d Cir. 1987). With the Drug Price
14 Competition and Patent Term Restoration Act of 1984, Congress relaxed the procedure for obtaining

15 approval from the FDA to market and sell a generic drug, allowing the generic maker to submit an
16 abbreviated NDA (“ANDA”). Id.; see 21 U.S.C. § 355(j), 35 U.S.C. §§ 156, 271, 281.

17 The ANDA must certify that the generic manufacturer will produce a bio-equivalent of the listed
18 drug and that the labeling and warnings of the generic drug are the same as those of the listed drug.
19 21 U.S.C. § 355(j)(2)(A).

20 **C. Procedural History**

21 On October 12, 2005, Plaintiff Margarita Gaeta filed a Complaint, as guardian ad litem for
22 her son, A.G., against Perrigo and Longs Drug Stores Corporation. (Complaint, Docket Item No. 1.)

23 The Complaint has been amended twice, adding Augustine Gaeta as a Plaintiff and PAR and BASF
24 as Defendants. (See Docket Item Nos. 6, 29.) On February 28, 2006, Plaintiffs filed a Second
25 Amended Complaint, which remains the operative complaint. In the Second Amended Complaint,

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1 Plaintiffs allege that Defendants are liable for injuries A.G. sustained as a result of ingesting
2 ibuprofen manufactured and distributed by Defendants. (Id.)

3 Plaintiffs allege the following causes of action against Defendants: (1) Defective Design; (2)

4 Marketing Defect; (3) Breach of Express Warranty; (4) Breach of Implied Warranty; (5) Negligence
5 and Gross Negligence; and (6) Deceit by Concealment pursuant to Cal. Civ. Code §§ 1709-1710.
6 (SAC ¶¶ 33-53.)

7 Presently before the Court is Perrigo's motion for summary judgment.

8 **III. STANDARDS**

9 Summary judgment is proper "if the pleadings, depositions, answers to interrogatories, and
10 admissions on file, together with the affidavits, if any, show that there is no genuine issue as to any
11 material fact and that the moving party is entitled to judgment as a matter of law." Fed. R. Civ. P.
12 56(c). The purpose of summary judgment "is to isolate and dispose of factually unsupported claims
13 or defenses." *Celotex v. Catrett*, 477 U.S. 317, 323-24 (1986).

14 The moving party "always bears the initial responsibility of informing the district court of
15 the basis for its motion, and identifying the evidence which it believes demonstrates the absence of a
16 genuine issue of material fact." *Id.* at 323. The non-moving party must then identify specific facts
17 "that might affect the outcome of the suit under the governing law," thus establishing that there is a
18 genuine issue for trial. Fed. R. Civ. P. 56(e).

19 When evaluating a motion for summary judgment, the court views the evidence through the

20prism of the evidentiary standard of proof that would pertain at trial. *Anderson v. Liberty Lobby*
21Inc., 477 U.S. 242, 255 (1986). The court draws all reasonable inferences in favor of the non-
22moving party, including questions of credibility and of the weight that particular evidence is
23accorded. See, e.g., *Masson v. New Yorker Magazine, Inc.*, 501 U.S. 496, 520 (1992). The court
24determines whether the non-moving party’s “specific facts,” coupled with disputed background or
25contextual facts, are such that a reasonable jury might return a verdict for the non-moving party.
26*T.W. Elec. Serv. v. Pac. Elec. Contractors*, 809 F.2d 626, 631 (9th Cir. 1987). In such a case,

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1summary judgment is inappropriate. *Anderson*, 477 U.S. at 248. However, where a rational trier of
2fact could not find for the non-moving party based on the record as a whole, there is no “genuine
3issue for trial.” *Matsushita Elec. Indus. Co. v. Zenith Radio*, 475 U.S. 574, 587 (1986).

4**IV. DISCUSSION**

5Perrigo moves for summary judgment on the ground that all of Plaintiffs’ claims are
6preempted by the FDCA. (Motion at 7.)

7The Supremacy Clause of United States Constitution provides that federal laws and treaties

8 “shall be the supreme law of the land.” U.S. Const. Art. VI, Cl. 2. The United States Supreme
9 Court has recognized three types of federal preemption of state law under the Supremacy Clause: (1)
10 express preemption, where Congress states explicitly the preemptive effect of its legislation on state
11 law; (2) field preemption, where Congress intends for federal law to occupy exclusively an entire
12 field of regulation; and (3) conflicts preemption, where it is impossible for a private party to comply
13 with both state and federal requirements. *English v. General Electric Co.*, 496 U.S. 72, 78-79
14 (1990). 15 The Food and Drug Modernization Act of 1997 (“Modernization Act”) amended the
FDCA
16 to provide for express preemption of state laws regarding non-prescription, “OTC” drugs. 21 U.S.C.
17 § 379r(a). However, § 379 has a savings clause, which provides that preemption provision does not
18 affect “the liability of any person under the product liability law of any State.” § 379r(e). The scope
19 of the term “product liability law” as used in the statute is not exactly clear. The California Court of
20 Appeal has found that the § 379r savings clause does not cover all “common law and statutory
21 actions imposing liability on commercial sellers of products.” *Kanter v. Warner-Lambert Co.*, 99
22 Cal. App. 4th 780, 790-91 (2002). Regardless, the savings clause “does not bar the ordinary
23 working of conflict pre-emption principles.” *Geier v. Am. Honda Motor Co., Inc.*, 529 U.S. 861,

24869 (2000). Under such principles, a court should “decline to give broad effect to savings clauses
25 where doing so would upset the careful regulatory scheme established by federal law.” United

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1 States v. Locke, 529 U.S. 89, 106-07 (2000). Thus, the Court proceeds to consider whether conflicts
2 between common law and federal law upset the regulatory scheme of the FDA.

3 Consideration of conflicts preemption under the Supremacy Clause “starts with the basic
4 assumption that Congress did not intend to displace state law.” Building and Const. Trades Council
5 of Metro. Dist. v. Assoc. Builders and Contractors of Mass./RI, Inc., 507 U.S. 218, 224 (1993). In
6 order for a court to find conflicts preemptions, there must be “clear evidence of a conflict.” Geier,
7 529 U.S. at 885. Conflicts preemption can occur with respect to the regulations of a federal agency
8 because regulations promulgated pursuant to federal statutory authority “have no less pre-emptive
9 effect than federal statutes.” Fidelity Federal Savings and Loan Ass’n v. de la Cuesta, 458 U.S. 141,
10 153 (1982). The preemption of such regulations may be challenged only to determine whether they
11 exceed statutory authority or were made arbitrarily. Id.

12 In approving an ANDA for a generic drug, the FDA requires the drug’s manufacturer “to

13 show that the labeling proposed for the drug is the same as the labeling approved for the listed drug
14 referred to in the [ANDA].” 21 C.F.R. § 314.127; 21 U.S.C. § 355(j)(2)(A)(v). “Labeling” is
15 defined by statute as “all labels and other written, printed, or graphic matter (1) upon any article or
16 any of its containers or wrappers, or (2) accompanying such article.” 21 U.S.C § 321(m). Thus,
17 labeling “embraces advertising or descriptive matter that goes with the package in which the articles
18 are transported,” in addition to any label that may be placed directly on a pill bottle. *Kordel v.*
19 *United States*, 335 U.S. 345, 350 (1948). The FDA will allow certain changes to the label in an
20 approved petition under § 314.93, which provides:

21 22 23^A is which combination application. not person identical one who active drug, wants to
ingredient a must listed to submit first drug is obtain substituted in an route abbreviated permission of
administration, for one new from of drug the FDA active application dosage to submit ingredients form,
for such and a drug an in strength, abbreviated a product listed or which in 24 21 C.F.R. § 314.93.

25 The FDA may withdraw approval for an ANDA if the agency finds that “the labeling for the
26 drug product . . . is no longer consistent with that for the listed drug” or that the label “is false or

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of a causal association with the drug.” 73 Fed. Reg.
le). However, “CBE changes are not available for
. . . . To the contrary, a generic drug manufacturer is
g for the listed drug.” Id.; see 21 CFR 314.150(b)(10);
der these regulations, a generic drug manufacturer
contraindication without FDA approval.²
ations and set forth its position that “under existing
eeling under the act . . . preempts conflicting or contrary
DA’s reasons for this position are as follows:
opagate interpretations of the act and FDA regulations
interpretations and frustrate the agency’s implementation
le, courts have rejected preemption in State law
that a manufacturer has latitude under FDA regulations
gthening warning statements without first obtaining
etermination whether labeling revisions are necessary is,
’s under the act. . . .
v serves as an appropriate source of supplementary safety
; or requiring manufacturers to disseminate risk

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7 FDA under the act. In fact, FDA interprets the act to
1g,” such that additional disclosures of risk information
ty under the act if the additional statement is
r misleading. Given the comprehensiveness of FDA
ess, and labeling under the act, additional requirements

C.F.R. § 314.150(b)(3), (b)(10). For a non-generic drug, a

) supplement to a label “is appropriate to amend the labeling for an

strengthen a contraindication, warning, precaution, or adverse

information are not necessarily more protective of patients. Instead, of an indication or other aspect of labeling protected
not the careful and truthful representation of benefits and risks that
appropriate judgments about drug use. Exaggeration of risk could 7
use of a beneficial drug. . . .

generic drug may only unilaterally change its label to reflect

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¹234^{FDA} in Overwarning, and labeling safe undermining scientific public and ^{has} that effective
previously health. evidence does the just objectives not use like Similarly, ^{found} can accurately of
underwarning, approved cause ^{that} of State-law the ^{labeling} meaningful portray act. products can ^{that}
attempts a similarly product's or risk ^{includes} encouraging information to have impose risks,
theoretical ^a thereby negative inappropriate additional to "lose ^{hazards} potentially effect its warnings
significance." ^{not} use on well-grounded patient and discouraging can lead safety to ⁵Id. at 3934-35
(citations ommitted).

⁶In the absence of clear authority to the contrary, a court is to give deference to an agency's

⁷interpretation of the scope of its authority to regulate. *Chevron U.S.A., Inc. v. Natural Resources*

⁸*Defense Council, Inc.*, 467 U.S. 837, 843-44 (1984); see *Medtronic, Inc. v. Lohr*, 518 U.S. 470, 505-

⁹06 (1996) (Breyer, J., concurring) (considering preemptive effect of FDA regulations, in light of the

¹⁰FDA's position that certain claims were not preempted). Under this principle, "state tort law which

11 would hold a generic drug manufacturer liable for failing to modify a label . . . conflict[s] with the
12 FDCA,” and any such claims are preempted by FDA regulations to the extent they seek to do so.
13 Colacicco v. Apotex, Inc., 432 F. Supp. 2d 514, 537-38 (E.D. Pa. 2006) aff’d, Colacicco v. Apotex
14 Inc., 2008 WL 927848 (3d Cir. 2008); cf. Papike v. Tambrands Inc., 107 F.3d 737, 742 (9th Cir.
15 1997).

16 In this case, Plaintiffs allege causes of action for: (1) Defective Design; (2) Marketing
17 Defect; (3) Breach of Express Warranty; (4) Breach of Implied Warranty; (5) Negligence and Gross
18 Negligence; and (6) Deceit by Concealment. (SAC ¶¶ 33-53.) With respect to each of these causes
19 of action, Plaintiffs allege, at least in part, that Defendants failed to warn individuals with
20 appropriate materials. (Id.) Specifically, in pleading their Defective Design cause of action,
21 Plaintiffs allege that Perrigo “failed to adequately and completely inform or warn of the risks of liver
22 injury and renal failure associated with the use of OTC ibuprofen to treat children for pain or fever.”
23 (Id. ¶ 35.) In their Marketing Defect cause of action, Plaintiffs allege that “[t]he warnings and
24 instructions that accompanied the defendants’ drugs provided inadequate warning to the consumer
25 and/or healthcare provider about the risk of acute liver failure.” (Id. ¶ 40.) Similar allegations are
26 made in Plaintiffs’ Breach of Warranty, Negligence, and Deceit causes of action. (Id. ¶¶ 42-53.)

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ved by the FDA under the ANDA process in 1987, which
ective. (Declaration of Robert Pinco in Support of
er, “Pinco Decl.,” Docket Item No. 159.) In 2002, the
egarding the safety of ibuprofen. (Id. ¶ 8.) The FDA
iry was not scientifically supported by the available data.
lso considered warning for risk of kidney injury and
ibuprofen should have a warning directed [to] those at
re associated with the use of the product.” Id. at 54144-
d inclusion of the warning. 71 Fed. Reg. 77314, 77316.
.G., Perrigo’s ibuprofen followed the labeling for the
nandated by the FDA. (Pinco Decl. ¶¶ 7, 8.) Thus, the
the labeling requirements that the FDA has set for OTC
to hold Perrigo liable for, in part, failing to warn of risks
; these warning would put the Perrigo’s ANDA in
A’s approved labeling for the listed drug, Plaintiffs’ state
obligations under federal law.

1 part Perrigo’s Motion for Summary Judgment. The

s of action are preempted to the extent that they allow for liability

ning on the company's OTC generic drug labeling for its 200mg

V. CONCLUSION

o moved for an extension of time to respond to

rriigo's Motion for Summary Judgment. Perrigo's Motion to Strike
, 225.)9

tt, an expert designated to testify about warning labels, is DENIED

The Court defers entering judgment in favor of Perrigo. The Court sets a Further Case

Management Conference for **June 30, 2008 at 10 a.m.** The parties shall meet and confer and file a

Joint Case Management Statement on or before **June 20, 2008.** The Statement shall address what

claims, if any, remain at issue in this case. Specifically, whether any of Plaintiffs' claims are based

on another non-preempted theory of recovery, such as, design defect for failure to conform to the

specification of the FDA approved form of the drug.

This Order terminates Docket Item Nos. 156, 188, 221, and 225.

Dated: June 13, 2008 JAMES United States WARE District Judge

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ing, Clerk

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By: /s/ JW Chambers

Elizabeth Garcia

Courtroom Deputy

Colin C. Munro

leen T. Davies

Kay Dopson