



U.S. Illnesses and Deaths Associated With Compounded or Repackaged Medications, 2001-17

Update note: This chart was updated in February 2019 to include newly reported adverse events, remove previously listed events following additional investigation, and update information in citations.

Pew’s drug safety project has identified 69 reported compounding errors or potential errors associated with more than 1,418 adverse events, including 114 deaths, from 2001 to 2017. However, a 2015 survey found that only 30 percent of states (13 of the 43 that responded) require sterile compounding pharmacies to report serious adverse events.¹ Of the states that require reporting, the type of information that is required to be reported may vary, further contributing to an incomplete picture of adverse events associated with compounded medications. Even in states with strong adverse event reporting requirements, illnesses and deaths caused by compounded drugs are not always linked to the compounding error.² Because many such events go unreported, this chart is an underestimation of the number of compounding errors since 2001. Contamination of sterile products was the most common error; others were the result of compounders’ miscalculations and mistakes in filling prescriptions.

Drug compounding can be an interstate operation; compounders may prepare medicines in one state and ship them to another. States may encounter oversight challenges if an out-of-state compounder shipping into their jurisdiction is held to a different quality or regulatory standard than in-state compounders. As a result, for each row below, the state where the compounding error or potential error occurred and the state(s) where the adverse event(s) occurred are listed. Harmonized minimum quality standards for anyone who compounds drugs—in any setting—across states would help address challenges in regulating out-of-state compounders and ensure that all compounding meets strong baseline criteria for preparing safe drugs and protecting patients.

Year	Reported cases	Reported deaths	Adverse event(s)	Compounding error	Product	State where compounding error occurred	State(s) where adverse event(s) occurred	Notes
2017 ³	At least 43		Vision impairment, poor night vision, loss of color perception, photophobia, ocular discomfort, nausea, loss of balance ⁴	Product contained multiple substances, including poloxamer 407 and poloxamer 407 degradants	Injectable steroid antibiotic combination for administration in the eye	TX	TX	

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2017 ⁵	2	1	One case of cardiac arrest; both experienced immediate hypersensitivity reactions	Product contained ungraded PEG 40 castor oil	Injectable curcumin emulsion infusion	CA	Not reported	
2017 ⁶	41		Septic arthritis	Bacterial contamination	Intra-articular injectable	NJ	NJ	Investigation revealed inappropriate use and handling of pharmacy bulk packaged products.
2017 ⁷	1		Hemorrhagic occlusive retinal vasculitis	Not reported	Intraocular injectable of triamcinolone, moxifloxacin, and vancomycin (TMV)	NJ	Not reported	
2017 ⁸	2		Tissue erosion at injection site	High pH; no glutamine detected in samples	Compounded injectable of glutamine, arginine, and carnitine (GAC)	FL	Not reported	
2016 ⁹	17	2	Fungal bloodstream infections	Contamination	Injectable saline, heparin, vancomycin, and ceftazidime	NY	NY	IV flush solutions were not compounded under quality standards set by the United States Pharmacopeial Convention and were used past appropriate beyond-use dating. The two deaths occurred within 12 weeks of the fungal infection, but it is unclear whether the deaths were a result of the infections.
2016 ¹⁰	1		Overdose	Dose of manganese chloride 1,000 times stronger than usual dose	Injectable manganese chloride	Not reported	Not reported	High manganese dose of 800 mg, compared with usual dose of 0.15-0.8 mg/day. Patient showed no resulting symptoms, but manganese overdose can result in side effects on the nerves and brain.
2016 ¹¹	3		Unspecified serious adverse events	Dose of morphine sulfate stronger than labeled concentration	Injectable morphine sulfate	IN	IL, IN ¹²	

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2016 ¹³	1		Abscesses and osteomyelitis	Contamination	Unknown injectable	Not reported	NM	Investigation revealed unsafe injection and compounding practices.
2016 ¹⁴	7		Thyrotoxicosis	Super-potent compounded drug	Compounded oral liothyronine	SD	Not reported	
2015 ¹⁵	7		Hepatitis C	Contamination	Unknown injectable	CA	CA	Investigation into the clinic revealed infection control breaches and ongoing issues with infection control practices.
2015 ¹⁶	Several		Unspecified	Adulterated and misbranded drug product (contained different API)	L-citrulline	NY	Not reported	Some samples of the product were found to contain a different amino acid (N-Acetyl-Leucine) than what the label claimed, and others did not contain any L-Citrulline.
2015 ¹⁷	5		Redness, swelling, and pain at injection site	Contamination	Compounded betamethasone phosphate and betamethasone acetate	AL	Not reported	
2015 ¹⁸	"Several" ¹⁹		Unspecified	High dose of vitamin D3	Oral multivitamin capsule	FL	Nationwide ²⁰	High vitamin D ₃ can cause significant short- and long-term effects.
2014-15 ²¹	"Several" ²²		Unspecified	Contamination	Sterile products	AL	Nationwide ²³	Administration of contaminated sterile products may result in serious and potentially life-threatening infections or death.
2014 ²⁴	Unknown		Oversedation	Dose of midazolam labeled with incorrect concentration	Injectable midazolam	IN	Not reported	Compounded midazolam, a sedating agent, did not match the concentration on the product label. Oversedation can result in a range of effects from increased sleepiness to severe difficulty breathing.

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2014 ²⁵	1	1	Toxicity	Not reported	Compounded topical anesthetic cream (ketamine)	TX	TX	
2014 ²⁶	37		Not reported	Contamination	Intravitreal injections of bevacizumab or ranibizumab	FL	Not reported	Bevacizumab and ranibizumab were repackaged in a manner that exposed sterile, preservative free vials to an uncontrolled environment.
2014 ²⁷	1		Severe flushing, stinging, and dizziness	Dose of magnesium sulfate 200 times stronger than labeled concentration ²⁸	Compounded magnesium sulfate	TX	Not reported	
2013 ²⁹	1		Bacterial bloodstream infection	Contamination	Injectable mineral product	TX	CA	Voluntary recall of injectable mineral product that contained bacteria with the potential for serious infection. A patient admitted to the hospital with an infection of the same bacteria.
2013 ³⁰	15	2 ³¹	Bacterial bloodstream infection	Contamination	Injectable calcium gluconate	TX	TX	The Centers for Disease Control and Prevention (CDC) has not conclusively linked the deaths to the contaminated drug.
2013 ³²	6		Fever, flu-like symptoms, soreness at injection site	Unknown	Injectable methylcobalamin	TX	Not reported	A compounded injection was recalled due to complaints of fever, flu-like symptoms, and soreness at the injection site. Subsequent Food and Drug Administration inspection found that sterility and quality of the manufacturing process could not be assured.
2013 ³³	5		Serious bacterial eye infections	Contamination	Injectable bevacizumab for administration in the eye	GA	GA, IN	

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2013 ³⁴	8		Fungal eye infections	Contamination	Injectable bevacizumab-triamcinolone for administration in the eye	Not reported	NY	Fungal infection of the eye caused significant visual impairment that persisted for at least three months from the incident.
2013 ³⁵	1		Kidney failure and acute injury of the liver and pancreas	Unknown	Injectable combination product for administration under the skin	Not reported	Not reported	Product is marketed for dissolving fat. The patient developed difficulties with digestion and metabolism as well as kidney failure, which required dialysis.
2012-13 ³⁶	12		Bacterial bloodstream infection	Contamination	Parenteral infusion	Not reported	IL	Facility inspection revealed deficiencies in the parenteral medication preparation and handling.
2012-13 ³⁷	26		Bacterial and fungal infections in skin and soft tissue	Contamination	Injectable preservative-free methylprednisolone acetate	TN	AR, FL, IL, NC	Skin and soft tissue infections resulted after intramuscular injection of preservative-free product. Subsequent voluntary recall of sterile products was issued.
2012-13 ³⁸	793	76 ³⁹	Fungal meningitis and other infections	Contamination	Injectable preservative-free methylprednisolone acetate	MA	FL, GA, ID, IL, IN, MD, MI, MN, NC, NH, NJ, NY, OH, PA, RI, SC, TN, TX, VA, WV ⁴⁰	Additional products (betamethasone, cardioplegia, and triamcinolone solutions) produced at the facility were also found to be contaminated, but adverse events linked to these products have not been reported. ⁴¹
2012 ⁴²	47		Fungal eye infection; vision loss in majority of cases	Contamination	Injectable brilliant blue-G (BBG) retinal dye and triamcinolone for administration in the eye	FL	CA, CO, IL, IN, LA, NC, NV, NY, TX	
2012 ⁴³	7		Bacterial bloodstream infection	Contamination	Injectable fentanyl	NC	NC	

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2012 ⁴⁴	1		Overdose	Dose of flecainide four times stronger than ordered	Oral flecainide liquid	Not reported	Not reported	Flecainide toxicity can cause abnormal heart rate and rhythms that can be severe and life-threatening, as well as increased liver enzymes, which can be an indicator of liver injury.
2012 ⁴⁵	10	1	Bacterial bloodstream infection	Contamination	Contrast dye, anesthetic and steroid injections-single-dose vials	Not reported	AZ, DE	The outpatient pain clinic failed to follow Standard Precautions by using single-dose vials as multi-dose vials. ⁴⁶
2011-12 ⁴⁷	15		Bacterial bloodstream infection	Contamination	Sterile products	Not reported	WV	Adverse events resulted from the use of bulk saline bag for IV flushes in a physician office practice.
2011 ⁴⁸	1		Toxicity	Dose of 4-aminopyridine 10 times stronger than labeled concentration	Oral 4-aminopyridine pills	Not reported	Not reported	Patient experienced stomach pain, anxiety, extreme sweating, and slow heart rate prior to developing life-threatening seizures. Following a complicated hospital stay, the patient sustained permanent short-term memory loss.
2011 ⁴⁹	9		Bacterial eye infection, and one case of meningitis and encephalitis; four cases of loss of eyesight	Contamination	Injectable bevacizumab for administration in the eye	Not reported	TN	
2011 ⁵⁰	12		Bacterial eye infection; three patients had eye removals	Contamination	Injectable bevacizumab for administration in the eye	FL	FL	
2011 ⁵¹	5		Blindness	Unintended presence of another medication	Injectable bevacizumab for administration in the eye	CA	CA	Trace amounts of bortezomib, a cancer drug that is not intended for injection into the eye, were detected on a sample syringe.

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2011 ⁵²	19	9	Bacterial bloodstream infection	Contamination	Parenteral nutrition solution	Not reported	AL	
2010 ⁵³	1	1	Fatal overdose	Dose of sodium 60 times stronger than ordered	Injectable sodium chloride	IL	IL	Acute sodium overload can cause symptoms ranging from fluid retention to seizures and coma, affecting multiple organs including lungs and kidneys. The patient was exposed to a potent sodium-containing fluid that was entered incorrectly during the preparation of the medication, resulting in death.
2010 ⁵⁴	1		Unspecified side effects	Dose of liothyronine 10 times stronger than ordered	Oral liothyronine (T3)	AZ	Not reported	Liothyronine overdose can result in shakiness, increased heart rate, and palpitations.
2009 ⁵⁵	1	1	Fatality	Unknown	Injectable hydromorphone	TN	Not reported	
2009 ⁵⁶	1	1	Fatal overdose	Dose of levothyroxine 18 times stronger than ordered	Oral levothyroxine pills	NC	Not reported	
2009 ⁵⁷	9		Eye infection; at least one case of vision loss	Unknown	Injectable preservative-free hyaluronidase for administration in the eye	FL	Not reported	Patients developed orbital cellulitis, a type of infection that results in inflammation of the eye.
2008 ⁵⁸	1		Acute withdrawal	Dose of baclofen 7 percent of ordered dosage	Injectable baclofen for administration in the spine	Not reported	Not reported	The product included a fraction of the intended dose of baclofen. Patient experienced frequent and severe spasms.
2008 ⁵⁹	1	1	Fatal overdose	Dose of sodium chloride 10 times stronger than ordered	Injectable sodium chloride	NC	Not reported	Acute sodium overload can cause symptoms ranging from fluid retention to seizures and coma, and affect multiple organs including lungs and kidneys.

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2008 ⁶⁰	1		Persistent inflammatory reaction	Unknown	Mesotherapy injections	Not reported	CO	Seven months after receiving mesotherapy injections, patient developed a persistent immune-mediated inflammatory reaction.
2007 ⁶¹	1	1	Fatal acute respiratory distress syndrome	Colistimethate sodium left in solution longer than recommended	Colistimethate sodium inhaled solution	Not reported	Not reported	The prodrug of colistin is better tolerated than the active drug to which it converts. More than half of the prodrug is converted within two days in solution at a certain temperature. This premixed product was in solution for five weeks before further dilution for administration.
2007 ⁶²	3	3	Fatal overdose	Dose of colchicine eight times stronger than labeled concentration	Injectable colchicine	TX	OR, WA	IV doses that exceed the standard single-use therapeutic dose of 2-4 mg per episode of gout have resulted in life-threatening toxicity. In this case, the doses were eightfold these limits.
2007 ⁶³	8	1	Bacterial bloodstream infection	Contamination	Injectable fentanyl	Not reported	CA, MD	
2006 ⁶⁴	1		Decreased consciousness, low blood pressure, and lack of oxygen	Mislabeled product leading to administration of different drug than ordered	Epidural morphine sulfate (fentanyl/ bupivacaine was ordered)	MS	AZ	Both fentanyl and morphine are in the same class of sedative analgesics. The symptoms of decreased consciousness, hypoxia, and hypotension are consistent with higher than intended opioid exposure.
2006 ⁶⁵	At least 70		Redness, swelling, bruising, rash, fever, and cellulitis	Betamethasone made with incorrect amount of preservative	Injectable betamethasone	AL	Not reported	The product was voluntarily recalled, and a subsequent reformulation continued to include an incorrect amount of preservative. An FDA investigation discovered at least 70 complaints associated with the drug.

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2006 ⁶⁶	1	1	Fatal overdose	Dose of chemotherapy infusion diluted with toxic amount of sodium chloride	Chemotherapy infusion	OH	OH	Acute sodium overload can cause symptoms ranging from fluid retention to seizures and coma, and affect multiple organs including lungs and kidneys.
2006 ⁶⁷	1	1	Fatal overdose	Dose of zinc 1,000 times stronger than ordered	Neonatal parenteral nutrition solution	NV	NV	The dose was incorrectly entered for pharmacy preparation as milligrams instead of micrograms, resulting in a thousandfold overdose.
2005 ⁶⁸	3	1	Fatal overdose, cardiac arrest	Dose of lidocaine and tetracaine higher than usual	Topical combination anesthetic creams (lidocaine and tetracaine)	NC	NC	
2005 ⁶⁹	19	1	Bacterial bloodstream infection	Contamination	Injectable magnesium sulfate	TX	CA, MA, NC, NJ, NY, SD	
2004-06 ⁷⁰	80		Bacterial bloodstream infection	Contamination	Injectable heparinized saline	TX	MI, MO, NY, SD, TX, WY	
2004-05 ⁷¹	6		Bacterial eye infection; all cases had partial or complete loss of vision; two patients had eye removals	Contamination	Trypan blue eye drops	Not reported	Not reported	

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2004-05 ⁷²	11	3	Systemic inflammatory response syndrome	Contamination	Cardioplegia solution for administration during heart surgery	MD	VA	
2004 ⁷³	2		Bacterial bloodstream infection	Contamination	Injectable heparin-vancomycin	FL	CT	
2003 ⁷⁴	2		Overdose	Dose of liothyronine stronger than ordered	Oral liothyronine (T3) pills	AZ	Not reported	Unused pills of both patients were analyzed, and the concentration of the active ingredient was found to be 800 and 900 times higher than intended. High T3 levels can result in shakiness, increased heart rate and palpitations.
2002-04 ⁷⁵	1	1	Fatal overdose	Dose of lidocaine and tetracaine higher than usual	Topical combination anesthetic cream (lidocaine and tetracaine)	UT	AZ	
2002 ⁷⁶	1		Toxicity	Dose of clonidine 10 times higher than ordered	Oral clonidine capsules	Not reported	Not reported	Patient showed early signs of central nervous system depression (somnia and drowsiness) and miosis (constricted or small pupils).
2002 ⁷⁷	1		Toxicity	Dose of clonidine 87 times higher than ordered	Oral clonidine liquid	Not reported	Not reported	Patient showed signs of central nervous system depression, consistent with severe clonidine toxicity. Miosis (constricted or small pupils) was also noted.

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2002 ⁷⁸	2		Meningitis	Contamination	Injectable methylprednisolone for administration in the spine	MI	MI	
2002 ⁷⁹	7	2	Fungal meningitis and sacroiliitis	Contamination	Injectable methylprednisolone acetate for administration in the spine	SC	NC	
2001 ⁸⁰	2		Bacterial bloodstream infection	Contamination	Injectable preservative-free heparinized saline	Not reported	Not reported	
2001 ⁸¹	1		Overdose	Dose of clonidine 1,000 times stronger than ordered	Oral clonidine liquid	Not reported	Not reported	During preparation of liquid clonidine from solid pills, milligrams were substituted for micrograms, resulting in a thousandfold overdose. Patient's initial presentation included hyperventilation, an unusual feature of clonidine toxicity. Severe clonidine toxicity can result in low blood pressure, central nervous system depression (lethargy, mental status changes), and cardiopulmonary instability (heart and breathing problems).
2001 ⁸²	13	3	Five cases of meningitis; five cases of epidural abscess; one patient had an infected hip joint; two unspecified	Contamination	Injectable betamethasone for administration in spine or joint	CA	CA	

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2001 ⁸³	4		Bacterial bloodstream infection	Contamination	Injectable ranitidine	Not reported	Not reported	

This chart includes U.S. illnesses and deaths associated with compounded or repackaged medications from 2001 to 2017. Adverse events were drawn from FDA and CDC resources as well as journal and news articles. In the total, “several” reported cases were counted as two adverse events, and an “unknown” number of reported cases were counted as zero adverse events.

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Endnotes

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