Register , 2018 <b>PROFESSIONAL REGULATIONS</b>
Chapter 52. Board of Pharmacy.
(Words in <b>boldface and underlined</b> indicate language being added; words [CAPITALIZED AND BRACKETED] indicate language being deleted. Complete new sections are not in boldface or underlined.)
12 AAC 52.855 is repealed and readopted to read:
12 AAC 52.855. Registration with the prescription drug monitoring program
controlled substance prescription database. (a) A licensed pharmacist shall register with the
Prescription Drug Monitoring Program's controlled substance prescription database (PDMP)
before dispensing a schedule II, III, or IV controlled substance under federal law.
(b) Before dispensing, prescribing, or administering a schedule II, III, or IV controlled
substance under federal law, a pharmacist or practitioner required to register with the PDMP
must
(1) register online on the PDMP website; and
(2) pay the fee established in 12 AAC 02.107.
(c) After completing the registration requirements, a pharmacist or practitioner required
to register with the PDMP will be issued a user account, login name, and password by the
department.
(d) A pharmacist or practitioner required to register with the PDMP must access
information in the PDMP using the user account, login name, and password issued by the
department.
(e) A pharmacist or practitioner required to register with the PDMP may access
information in the PDMP using another registrant's credentials only as authorized by a contract
executed by the department for the purposes of AS 47.07.038. (Eff. 12/29/2011, Register 200; am
/, Register)
<b>Authority:</b> AS 08.80.005 AS 08.80.030 AS 17.30.200

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12 AAC 52.860 is	repealed and readopted to read:
12 AAC 52	2.860. Access to and conditions for use of the prescription drug monitoring
program databas	se. (a) Access to the PDMP is limited as described in AS 17.30.200(d).
(b) For the	purposes of AS 17.30.200(d)(1):
(1)	"personnel of this board" means employees of the Department of Commerce,
Community, and I	Economic Development assigned to the Board of Pharmacy; and
(2)	"personnel of another board or agency" means an employee of the state of
Alaska assigned to	o a board that requires licensees to register with the PDMP or an agency
identified in a sear	rch warrant, subpoena, or order issued by an administrative law judge or a
court.	
(c) For the	purposes of AS 17.30.200(d)(2), "authorized board personnel or contractors"
means:	
(1)	employees of the Department of Commerce, Community, and Economic
Development assignment	gned to the Board of Pharmacy; or
(2)	employees of a state contractor providing PDMP data storage or data
management servi	ces.
(d) For the	purposes of AS 17.30.200(d)(3) and (4), a licensed practitioner or licensed or
registered pharma	cist authorizing an "agent or employee" to access the PDMP is responsible for
maintaining and te	erminating the agent or employee's access to the PDMP.
(e) For the	purposes of AS 17.30.200(d)(8) and (10), "authorized employee of the
Department of Hea	alth and Social Services" means an employee of the Department of Health and
Social Services (D	OHSS) for whom the DHSS commissioner or commissioner's official designee
has requested acce	ess in writing to the board prior to the release of information. (Eff. 12/29/2011,
Register 200; am _	/, Register)
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**Authority:** AS 08.80.005 AS 08.80.030 AS 17.30.200

12 AAC 52.865 is repealed and readopted to read:

**12 AAC 52.865. Reporting and reviewing PDMP information.** (a) Unless excused from reporting under AS 17.30.200(u), information required under AS 17.30.200(b) must be submitted by a pharmacist, if the pharmacist-in-charge is not present.

- (b) Unless excused from reporting under AS 17.30.200(u), a pharmacist or practitioner required to submit information under AS 17.30.200(b) must submit the information to the Alaska Prescription Drug Monitoring Program (PDMP) daily as of the previous submission date.
- (c) The time computation under 12 AAC 02.920(b) applies to a submission of information under AS 17.30.200(b) and this section.
- (d) For the purposes of AS 17.30.200(b)(8), "other appropriate identifier" and "other appropriate identifying information" means the state issued license number of the prescribing practitioner, and the dispensing pharmacist or practitioner.
- (e) Within 72 hours of discovering an error in information submitted under AS 17.30.200(b), a pharmacist or practitioner required to submit the information under AS 17.30.200(b) must submit information correcting the error to the PDMP administrator. The time computation under 12 AAC 02.920(b) applies to a submission of information correcting an error in information submitted under AS 17.30.200(b).
- (f) Unless excused from reporting under AS 17.30.200(u), or a waiver is granted under 12 AAC 52.870, a pharmacist or practitioner required to submit information under AS 17.30.200(b) must submit the information to the PDMP electronically through the website provided by the board.
  - (g) Unless excused from reviewing the PDMP under AS 17.30.200(k)(4)(A) (B), a

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practitioner, b	out not a pharmacist, m	oust review the inform	nation in the PDMP to	o check a patient's	
prescription re	ecords before dispension	ng, prescribing, or ad	ministering a schedul	e II or III	
controlled sub	ostance under federal la	aw. (Eff. 12/29/2011,	Register 200; am	_/,	
Register	_)				
Authority:	AS 08.80.005	AS 08.80.030	AS 17.30.200		
12 AAC 52.8°	70 is amended to read:				
12 AA	AC 52.870. Waiver of	electronic submissio	on requirement by <u>p</u>	harmacist or	
<u>practitioner</u>	[DISPENSER]. (a) Th	ne department shall w	aive the electronic su	bmission	
requirements	of <u>12 AAC 52.865(e)</u>	[12 AAC 52.865(b)]	for good cause. The <b>p</b>	harmacist or	
practitioner	[DISPENSER] request	ing the waiver is resp	onsible for establishi	ng the basis for the	
requested wai	ver under this section.				
(b) To	establish good cause t	for purposes of this se	ection, a <b>pharmacist</b>	or practitioner	
[DISPENSER	R] must submit an appl	ication and sworn sta	tement showing that		
(1) a natural disaster or other emergency beyond the control of the <b>pharmacist or</b>					
<u>practitioner</u> [DISPENSER] prevents the <u>pharmacist or practitioner</u> [DISPENSER] from					
complying wi	th <u>12 AAC 52.865(e)</u>	[12 AAC 52.865(b)];			
	(2) the <b>pharmacist o</b>	r practitioner [DISP	ENSER] will only di	spense controlled	
substances as	abstances as part of a controlled research project approved by an accredited institution of higher				
education or under the supervision of a government agency;					
	(3) repealed /	/[THE DISPEN	ISER WILL DISPEN	ISE NINE OR	
FEWER PRESCRIPTIONS OF CONTROLLED SUBSTANCES A MONTH];					
	(4) the <b>pharmacist's</b>	or practitioner's [D	ISPENSER'S] busine	ess is located in an	
area that lacks	s access to the telecom	munication services r	needed to comply with	h	

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<b>12 AAC 52.865(e)</b> [12 AAC 52.865(b)]; or
(5) the <b>pharmacist or practitioner</b> [DISPENSER] will suffer financial hardship
if required to acquire the technology necessary to comply with 12 AAC 52.865(e)
[12 AAC 52.865(b)].
(c) The department may not grant a waiver under this section unless the <b>pharmacist or</b>
<u>practitioner</u> [DISPENSER] first agrees in writing that, if the waiver is granted, the <u>pharmacist</u>
or practitioner [DISPENSER] will satisfy the reporting requirements of AS 17.30.200(b) by
submitting the required information by United States mail to the board on at least a daily basis
using a form approved by the board
[(1) THE PHARMACY UNIVERSAL CLAIMS FORM OF THE NATIONAL
COUNCIL FOR PRESCRIPTION DRUG PROGRAMS; OR
(2) AN ALTERNATIVE FORM APPROVED BY THE BOARD AS
PROVIDING SUBSTANTIALLY THE SAME INFORMATION AS THE FORM DESCRIBED
IN (1) OF THIS SUBSECTION].
(d) A request for a waiver under this section must be in writing using an application form
<b>provided</b> [PRESCRIBED] by the board and sent to the board.
(e) The department's grant or denial of a waiver request constitutes a final agency action
unless, no later than 30 days after the department issues notice of the grant or denial, the
pharmacist or practitioner [DISPENSER] files a written notice of appeal with the board.
(f) A waiver granted under this section expires at the end of the year in which it is
granted.
(g) A pharmacist or practitioner must [DISPENSER SHALL] inform the board within
30 days if the basis for the waiver of electronic reporting no longer exists. (Eff. 12/29/2011,
Register 200; am/, Register)
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Authority:	AS 08.80.005	AS 08.80.030	AS 17.30.200	
The introduct	ory language of 12	AAC 52.880(a) is amer	nded to read:	
(a) Th	e board will mainta	in a register for patient	profile requests solicited under	
[12 AAC 52.5	855(b) OR] 12 AAC	52.875. The register in	ncludes the following information:	
• • •				
12 AAC 52.8	80(a)(3) is amended	I to read:		
	(3) the name, title,	, [BUSINESS,] and add	lress of the individual requesting the	
profile [AND	, IF THE INDIVIDU	UAL IS A PRACTITIO	NER, THE PRACTITIONER'S	
CURRENT F	EDERAL DRUG E	ENFORCEMENT ADM	IINISTRATION REGISTRATION	
NUMBER];				
(Eff. 12/29/20	011, Register 200; a	m/, l	Register)	
Authority:	AS 08.80.005	AS 08.80.030	AS 17.30.200	
12 AAC 52 is	s amended by adding	g a new section to read:		
12 A	AC 52.885. Purge d	atabase records. The f	Collowing information will be purged	
from the PDN	MP database after tw	o years have elapsed fr	om the date the prescription was	
dispensed:				
	(1) the name of the	e prescribing practition	er and the practitioner's federal Drug	
Enforcement	Administration regi	stration number or othe	er appropriate identifier;	
	(2) the date of the	prescription;		
	(3) the date the pro	escription was filled and	d the method of payment;	
D 40/2/2=			f the person for whom the prescription	
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was written;					
	(5) the name and national drug code of the controlled substance;				
	(6) the quantity and strength of the controlled substance dispensed;				
	(7) the name of the drug outlet dispensing the controlled substance; and				
	(8) the name of the pharmacist or practitioner dispensing the controlled substance				
and other app	ropriate identifying in	formation. (Eff/_	/, Register)		
<b>Authority:</b>	AS 08.80.005	AS 08.80.030	AS 17.30.200		
12 AAC 52.89	90 is repealed and read	lopted to read:			
12 AA	AC 52.890. Grounds f	or discipline. A violat	ion of 12 AAC 52.855 –		
12 AAC 52.89	90 by a pharmacist is g	grounds for the imposit	ion of disciplinary sanctions under		
AS 08.01.075	and AS 08.80.261. A	violation of 12 AAC 5	22.855 – 12 AAC 52.890 by a		
practitioner no	ot licensed by this boa	rd shall be reported to	the practitioner's licensing board. (Eff.		
12/29/2011, F	Register 200; am/	/, Register	:)		
Authority:	AS 08.80.005	AS 08.80.030	AS 17.30.200		
12 AAC 52.9	20(a) is amended by ac	dding a new paragraph	to read:		
	(22) violating AS 17	.30.200 or a regulation	adopted thereunder dealing with the		
PDMP.					
(Eff. 1/16/98,	Register 145; am	_/, Regist	er)		
Authority:	AS 08.01.075	AS 08.80.261	AS 08.80.460		
	AS 08.80.005	AS 08.80.315	AS 17.30.200		
	AS 08.80.030				

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12 AAC 52.995 is amended by adding a new subsection to read:

(d) In AS 17.30.200, and 12 AAC 52.855 – 12 AAC 52.895, "practitioner" has the meaning given in AS 11.71.900. (Eff. 1/16/98, Register 145; am 5/5/2000, Register 154; am 11/10/2001, Register 160; am 8/21/2002, Register 163; am 2/15/2006, Register 177; am 8/12/2007, Register 183; am 9/11/2010, Register 195; am 12/29/2011, Register 200; am

8/1/2014, Register 211; am \_\_\_\_/\_\_\_, Register \_\_\_\_\_)

**Authority:** AS 08.80.005 AS 08.80.157 **AS 17.30.200** 

AS 08.80.030 **AS 11.71.900**