

**BEFORE THE
MEDICAL BOARD OF CALIFORNIA
DEPARTMENT OF CONSUMER AFFAIRS
STATE OF CALIFORNIA**

**In the Matter of the Accusation
Against:**

F. Omar Briones Tordilla, M.D.

**Physician's and Surgeon's
Certificate No. A 88578**

Case No.: 800-2022-086193

Respondent.

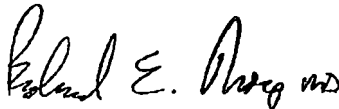
DECISION

The attached Stipulated Settlement is hereby adopted as the Decision and Order of the Medical Board of California, Department of Consumer Affairs, State of California.

This Decision shall become effective at 5:00 p.m. on December 1, 2025.

IT IS SO ORDERED: October 30, 2025.

MEDICAL BOARD OF CALIFORNIA



**Richard E. Thorp, M.D., Chair
Panel B**

1 ROB BONTA
Attorney General of California
2 MATTHEW M. DAVIS
Supervising Deputy Attorney General
3 ANDRES T. CARNAHAN
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7 *Attorneys for Complainant*

8
9 **BEFORE THE**
10 **MEDICAL BOARD OF CALIFORNIA**
11 **DEPARTMENT OF CONSUMER AFFAIRS**
12 **STATE OF CALIFORNIA**

13 In the Matter of the Accusation Against:

14 **F. OMAR BRIONES TORDILLA, M.D.**
15 **225 E Second Avenue, Suite 101**
16 **Escondido, CA 92025**

17 **Physician's and Surgeon's Certificate No. A**
18 **88578**

19 Respondent.

Case No. 800-2022-086193

OAH No. 2025040245

20 **STIPULATED SETTLEMENT AND**
21 **DISCIPLINARY ORDER**

22 IT IS HEREBY STIPULATED AND AGREED by and between the parties to the above-
23 entitled proceedings that the following matters are true:

24 **PARTIES**

25 1. Reji Varghese (Complainant) is the Executive Director of the Medical Board of
26 California (Board). He brought this action solely in his official capacity and is represented in this
27 matter by Rob Bonta, Attorney General of the State of California, by Andres T. Carnahan, Deputy
28 Attorney General.

2. Respondent F. Omar Briones Tordilla, M.D. (Respondent) is represented in this
proceeding by attorney Taryn A. Perez, Esq., whose address is: 527 Encinitas Blvd., Suite 100
Encinitas, California 92024.

1 3. On or about August 13, 2004, the Board issued Physician's and Surgeon's Certificate
2 No. A 88578 to F. Omar Briones Tordilla, M.D. (Respondent). The Physician's and Surgeon's
3 Certificate was in full force and effect at all times relevant to the charges brought in Accusation
4 No. 800-2022-086193, and will expire on November 30, 2025, unless renewed.

5 **JURISDICTION**

6 4. Accusation No. 800-2022-086193 was filed before the Board, and is currently
7 pending against Respondent. The Accusation and all other statutorily required documents were
8 properly served on Respondent on January 8, 2025. Respondent timely filed his Notice of
9 Defense contesting the Accusation.

10 5. A copy of Accusation No. 800-2022-086193 is attached as exhibit A and incorporated
11 herein by reference.

12 **ADVISEMENT AND WAIVERS**

13 6. Respondent has carefully read, fully discussed with counsel, and understands the
14 charges and allegations in Accusation No. 800-2022-086193. Respondent has also carefully read,
15 fully discussed with his counsel, and understands the effects of this Stipulated Settlement and
16 Disciplinary Order.

17 7. Respondent is fully aware of his legal rights in this matter, including the right to a
18 hearing on the charges and allegations in the Accusation; the right to confront and cross-examine
19 the witnesses against him; the right to present evidence and to testify on his own behalf; the right
20 to the issuance of subpoenas to compel the attendance of witnesses and the production of
21 documents; the right to reconsideration and court review of an adverse decision; and all other
22 rights accorded by the California Administrative Procedure Act and other applicable laws.

23 8. Respondent voluntarily, knowingly, and intelligently waives and gives up each and
24 every right set forth above.

25 **CULPABILITY**

26 9. Respondent admits the truth of each and every charge and allegation in Accusation
27 No. 800-2022-086193.

10. Respondent agrees that his Physician's and Surgeon's Certificate is subject to discipline and agrees to be bound by the Board's probationary terms as set forth in the Disciplinary Order below.

CONTINGENCY

11. This stipulation shall be subject to approval by the Medical Board of California. Respondent understands and agrees that counsel for Complainant and the staff of the Medical Board of California may communicate directly with the Board regarding this stipulation and settlement, without notice to or participation by Respondent or his counsel. By signing the stipulation, Respondent understands and agrees that he may not withdraw his agreement or seek to rescind the stipulation prior to the time the Board considers and acts upon it. If the Board fails to adopt this stipulation as its Decision and Order, the Stipulated Settlement and Disciplinary Order shall be of no force or effect, except for this paragraph, it shall be inadmissible in any legal action between the parties, and the Board shall not be disqualified from further action by having considered this matter.

12. This Stipulated Settlement and Disciplinary Order is intended by the parties herein to be an integrated writing representing the complete, final and exclusive embodiment of the agreement of the parties in this above entitled matter.

13. Respondent agrees that if he ever petitions for early termination or modification of probation, or if an accusation and/or petition to revoke probation is filed against him before the Board, all of the charges and allegations contained in Accusation No. 800-2022-086193 shall be deemed true, correct and fully admitted by respondent for purposes of any such proceeding or any other licensing proceeding involving Respondent in the State of California.

14. The parties understand and agree that Portable Document Format (PDF) and facsimile copies of this Stipulated Settlement and Disciplinary Order, including PDF and facsimile signatures thereto, shall have the same force and effect as the originals.

15. In consideration of the foregoing admissions and stipulations, the parties agree that the Board may, without further notice or opportunity to be heard by the Respondent, issue and enter the following Disciplinary Order:

1
2 **DISCIPLINARY ORDER**

3 IT IS HEREBY ORDERED that Physician's and Surgeon's Certificate No. A 88578 issued
4 to Respondent F. Omar Briones Tordilla, M.D. is revoked. However, the revocation is stayed and
5 Respondent is placed on probation for five (5) years on the following terms and conditions:

6 1. **CONTROLLED SUBSTANCES - TOTAL RESTRICTION.** Until such time as
7 Respondent has successfully completed the Clinical Competence Assessment Program described
8 in condition 7, below, Respondent shall not order, prescribe, dispense, administer, furnish, or
9 possess any controlled substances as defined in the California Uniform Controlled Substances
10 Act.

11 Until such time as Respondent has successfully completed the Clinical Competence
12 Assessment Program described in condition 7, below, Respondent shall not issue an oral or
13 written recommendation or approval to a patient or a patient's primary caregiver for the
14 possession or cultivation of marijuana for the personal medical purposes of the patient within the
15 meaning of Health and Safety Code section 11362.5.

16 If Respondent forms the medical opinion, after an appropriate prior examination and a
17 medical indication, that a patient's medical condition may benefit from the use of marijuana,
18 Respondent shall so inform the patient and shall refer the patient to another physician who,
19 following an appropriate prior examination and a medical indication, may independently issue a
20 medically appropriate recommendation or approval for the possession or cultivation of marijuana
21 for the personal medical purposes of the patient within the meaning of Health and Safety Code
22 section 11362.5. In addition, Respondent shall inform the patient or the patient's primary
23 caregiver that Respondent is prohibited from issuing a recommendation or approval for the
24 possession or cultivation of marijuana for the personal medical purposes of the patient and that
25 the patient or the patient's primary caregiver may not rely on Respondent's statements to legally
26 possess or cultivate marijuana for the personal medical purposes of the patient. Respondent shall
27 fully document in the patient's chart that the patient or the patient's primary caregiver was so
28 informed. Nothing in this condition prohibits Respondent from providing the patient or the

1 patient's primary caregiver information about the possible medical benefits resulting from the use
2 of marijuana.

3 2. CONTROLLED SUBSTANCES - MAINTAIN RECORDS AND ACCESS TO
4 RECORDS AND INVENTORIES. Respondent shall maintain a record of all controlled
5 substances ordered, prescribed, dispensed, administered, or possessed by Respondent, and any
6 recommendation or approval which enables a patient or patient's primary caregiver to possess or
7 cultivate marijuana for the personal medical purposes of the patient within the meaning of Health
8 and Safety Code section 11362.5, during probation, showing all of the following: 1) the name and
9 address of the patient; 2) the date; 3) the character and quantity of controlled substances involved;
10 and 4) the indications and diagnosis for which the controlled substances were furnished.

11 Respondent shall keep these records in a separate file or ledger, in chronological order. All
12 records and any inventories of controlled substances shall be available for immediate inspection
13 and copying on the premises by the Board or its designee at all times during business hours and
14 shall be retained for the entire term of probation.

15 3. EDUCATION COURSE. Within 60 calendar days of the effective date of this
16 Decision, and on an annual basis thereafter, Respondent shall submit to the Board or its designee
17 for its prior approval educational program(s) or course(s) which shall not be less than 40 hours
18 per year, for each year of probation. The educational program(s) or course(s) shall be aimed at
19 correcting any areas of deficient practice or knowledge and shall be Category I certified. The
20 educational program(s) or course(s) shall be at Respondent's expense and shall be in addition to
21 the Continuing Medical Education (CME) requirements for renewal of licensure. Following the
22 completion of each course, the Board or its designee may administer an examination to test
23 Respondent's knowledge of the course. Respondent shall provide proof of attendance for 65
24 hours of CME of which 40 hours were in satisfaction of this condition.

25 4. PRESCRIBING PRACTICES COURSE. Within 60 calendar days of the effective
26 date of this Decision, Respondent shall enroll in a course in prescribing practices approved in
27 advance by the Board or its designee. Respondent shall provide the approved course provider
28 with any information and documents that the approved course provider may deem pertinent.

Respondent shall participate in and successfully complete the classroom component of the course not later than six (6) months after Respondent's initial enrollment. Respondent shall successfully complete any other component of the course within one (1) year of enrollment. The prescribing practices course shall be at Respondent's expense and shall be in addition to the Continuing Medical Education (CME) requirements for renewal of licensure.

A prescribing practices course taken after the acts that gave rise to the charges in the Accusation, but prior to the effective date of the Decision may, in the sole discretion of the Board or its designee, be accepted towards the fulfillment of this condition if the course would have been approved by the Board or its designee had the course been taken after the effective date of this Decision.

Respondent shall submit a certification of successful completion to the Board or its designee not later than 15 calendar days after successfully completing the course, or not later than 15 calendar days after the effective date of the Decision, whichever is later.

5. MEDICAL RECORD KEEPING COURSE. Within 60 calendar days of the effective date of this Decision, Respondent shall enroll in a course in medical record keeping approved in advance by the Board or its designee. Respondent shall provide the approved course provider with any information and documents that the approved course provider may deem pertinent. Respondent shall participate in and successfully complete the classroom component of the course not later than six (6) months after Respondent's initial enrollment. Respondent shall successfully complete any other component of the course within one (1) year of enrollment. The medical record keeping course shall be at Respondent's expense and shall be in addition to the Continuing Medical Education (CME) requirements for renewal of licensure.

A medical record keeping course taken after the acts that gave rise to the charges in the Accusation, but prior to the effective date of the Decision may, in the sole discretion of the Board or its designee, be accepted towards the fulfillment of this condition if the course would have been approved by the Board or its designee had the course been taken after the effective date of this Decision.

Respondent shall submit a certification of successful completion to the Board or its

1 designee not later than 15 calendar days after successfully completing the course, or not later than
2 15 calendar days after the effective date of the Decision, whichever is later.

3 6. PROFESSIONALISM PROGRAM (ETHICS COURSE). Within 60 calendar days of
4 the effective date of this Decision, Respondent shall enroll in a professionalism program, that
5 meets the requirements of Title 16, California Code of Regulations (CCR) section 1358.1.
6 Respondent shall participate in and successfully complete that program. Respondent shall
7 provide any information and documents that the program may deem pertinent. Respondent shall
8 successfully complete the classroom component of the program not later than six (6) months after
9 Respondent's initial enrollment, and the longitudinal component of the program not later than the
10 time specified by the program, but no later than one (1) year after attending the classroom
11 component. The professionalism program shall be at Respondent's expense and shall be in
12 addition to the Continuing Medical Education (CME) requirements for renewal of licensure.

13 A professionalism program taken after the acts that gave rise to the charges in the
14 Accusation, but prior to the effective date of the Decision may, in the sole discretion of the Board
15 or its designee, be accepted towards the fulfillment of this condition if the program would have
16 been approved by the Board or its designee had the program been taken after the effective date of
17 this Decision.

18 Respondent shall submit a certification of successful completion to the Board or its
19 designee not later than 15 calendar days after successfully completing the program or not later
20 than 15 calendar days after the effective date of the Decision, whichever is later.

21 7. CLINICAL COMPETENCE ASSESSMENT PROGRAM. Within 60 calendar days
22 of the effective date of this Decision, Respondent shall enroll in a clinical competence assessment
23 program approved in advance by the Board or its designee. Respondent shall successfully
24 complete the program not later than six (6) months after Respondent's initial enrollment unless
25 the Board or its designee agrees in writing to an extension of that time.

26 The program shall consist of a comprehensive assessment of Respondent's physical and
27 mental health and the six general domains of clinical competence as defined by the Accreditation
28 Council on Graduate Medical Education and American Board of Medical Specialties pertaining to

1 Respondent's current or intended area of practice. The program shall take into account data
2 obtained from the pre-assessment, self-report forms and interview, and the Decision(s),
3 Accusation(s), and any other information that the Board or its designee deems relevant. The
4 program shall require Respondent's on-site participation as determined by the program for the
5 assessment and clinical education and evaluation. Respondent shall pay all expenses associated
6 with the clinical competence assessment program.

7 At the end of the evaluation, the program will submit a report to the Board or its designee
8 which unequivocally states whether the Respondent has demonstrated the ability to practice
9 safely and independently. Based on Respondent's performance on the clinical competence
10 assessment, the program will advise the Board or its designee of its recommendation(s) for the
11 scope and length of any additional educational or clinical training, evaluation or treatment for any
12 medical condition or psychological condition, or anything else affecting Respondent's practice of
13 medicine. Respondent shall comply with the program's recommendations.

14 Determination as to whether Respondent successfully completed the clinical competence
15 assessment program is solely within the program's jurisdiction.

16 If Respondent fails to enroll, participate in, or successfully complete the clinical
17 competence assessment program within the designated time period, Respondent shall receive a
18 notification from the Board or its designee to cease the practice of medicine within three (3)
19 calendar days after being so notified. The Respondent shall not resume the practice of medicine
20 until enrollment or participation in the outstanding portions of the clinical competence assessment
21 program have been completed. If the Respondent did not successfully complete the clinical
22 competence assessment program, the Respondent shall not resume the practice of medicine until a
23 final decision has been rendered on the accusation and/or a petition to revoke probation. The
24 cessation of practice shall not apply to the reduction of the probationary time period.

25 Within 60 days after Respondent has successfully completed the clinical competence
26 assessment program, Respondent shall participate in a professional enhancement program
27 approved in advance by the Board or its designee, which shall include quarterly chart review,
28 semi-annual practice assessment, and semi-annual review of professional growth and education.

Respondent shall participate in the professional enhancement program at Respondent's expense during the term of probation, or until the Board or its designee determines that further participation is no longer necessary.

8. MONITORING - PRACTICE Within 30 calendar days of the effective date of this Decision, Respondent shall submit to the Board or its designee for prior approval as a practice monitor, the name and qualifications of one or more licensed physicians and surgeons whose licenses are valid and in good standing, and who are preferably American Board of Medical Specialties (ABMS) certified. A monitor shall have no prior or current business or personal relationship with Respondent, or other relationship that could reasonably be expected to compromise the ability of the monitor to render fair and unbiased reports to the Board, including but not limited to any form of bartering, shall be in Respondent's field of practice, and must agree to serve as Respondent's monitor. Respondent shall pay all monitoring costs.

The Board or its designee shall provide the approved monitor with copies of the Decision(s) and Accusation(s), and a proposed monitoring plan. Within 15 calendar days of receipt of the Decision(s), Accusation(s), and proposed monitoring plan, the monitor shall submit a signed statement that the monitor has read the Decision(s) and Accusation(s), fully understands the role of a monitor, and agrees or disagrees with the proposed monitoring plan. If the monitor disagrees with the proposed monitoring plan, the monitor shall submit a revised monitoring plan with the signed statement for approval by the Board or its designee.

Within 60 calendar days of the effective date of this Decision, and continuing throughout probation, Respondent's practice shall be monitored by the approved monitor. Respondent shall make all records available for immediate inspection and copying on the premises by the monitor at all times during business hours and shall retain the records for the entire term of probation.

If Respondent fails to obtain approval of a monitor within 60 calendar days of the effective date of this Decision, Respondent shall receive a notification from the Board or its designee to cease the practice of medicine within three (3) calendar days after being so notified. Respondent shall cease the practice of medicine until a monitor is approved to provide monitoring responsibility.

1 The monitor(s) shall submit a quarterly written report to the Board or its designee which
2 includes an evaluation of Respondent's performance, indicating whether Respondent's practices
3 are within the standards of practice of medicine, and whether Respondent is practicing medicine
4 safely, billing appropriately or both. It shall be the sole responsibility of Respondent to ensure
5 that the monitor submits the quarterly written reports to the Board or its designee within 10
6 calendar days after the end of the preceding quarter.

7 If the monitor resigns or is no longer available, Respondent shall, within 5 calendar days of
8 such resignation or unavailability, submit to the Board or its designee, for prior approval, the
9 name and qualifications of a replacement monitor who will be assuming that responsibility within
10 15 calendar days. If Respondent fails to obtain approval of a replacement monitor within 60
11 calendar days of the resignation or unavailability of the monitor, Respondent shall receive a
12 notification from the Board or its designee to cease the practice of medicine within three (3)
13 calendar days after being so notified. Respondent shall cease the practice of medicine until a
14 replacement monitor is approved and assumes monitoring responsibility.

15 In lieu of a monitor, Respondent may participate in a professional enhancement program
16 approved in advance by the Board or its designee that includes, at minimum, quarterly chart
17 review, semi-annual practice assessment, and semi-annual review of professional growth and
18 education. Respondent shall participate in the professional enhancement program at Respondent's
19 expense during the term of probation.

20 9. SOLO PRACTICE PROHIBITION. Respondent is prohibited from engaging in the
21 solo practice of medicine. Prohibited solo practice includes, but is not limited to, a practice
22 where: 1) Respondent merely shares office space with another physician but is not affiliated for
23 purposes of providing patient care, or 2) Respondent is the sole physician practitioner at that
24 location.

25 If Respondent fails to establish a practice with another physician or secure employment in
26 an appropriate practice setting within 60 calendar days of the effective date of this Decision,
27 Respondent shall receive a notification from the Board or its designee to cease the practice of
28 medicine within three (3) calendar days after being so notified. The Respondent shall not resume

1 practice until an appropriate practice setting is established.

2 If, during the course of the probation, the Respondent's practice setting changes and the
3 Respondent is no longer practicing in a setting in compliance with this Decision, the Respondent
4 shall notify the Board or its designee within five (5) calendar days of the practice setting change.
5 If Respondent fails to establish a practice with another physician or secure employment in an
6 appropriate practice setting within 60 calendar days of the practice setting change, Respondent
7 shall receive a notification from the Board or its designee to cease the practice of medicine within
8 three (3) calendar days after being so notified. The Respondent shall not resume practice until an
9 appropriate practice setting is established.

10 10. NOTIFICATION. Within seven (7) days of the effective date of this Decision, the
11 Respondent shall provide a true copy of this Decision and Accusation to the Chief of Staff or the
12 Chief Executive Officer at every hospital where privileges or membership are extended to
13 Respondent, at any other facility where Respondent engages in the practice of medicine,
14 including all physician and locum tenens registries or other similar agencies, and to the Chief
15 Executive Officer at every insurance carrier which extends malpractice insurance coverage to
16 Respondent. Respondent shall submit proof of compliance to the Board or its designee within 15
17 calendar days.

18 This condition shall apply to any change(s) in hospitals, other facilities or insurance carrier.

19 11. OBEY ALL LAWS. Respondent shall obey all federal, state and local laws, all rules
20 governing the practice of medicine in California and remain in full compliance with any court
21 ordered criminal probation, payments, and other orders.

22 12. INVESTIGATION/ENFORCEMENT COST RECOVERY. Respondent is hereby
23 ordered to reimburse the Board its costs of investigation and enforcement, including, but not
24 limited to, expert review, amended accusations, legal reviews, investigation(s), and subpoena
25 enforcement, as applicable, in the amount of \$41,550.77 (forty-one thousand, five hundred fifty
26 dollars and seventy-seven cents). Costs shall be payable to the Medical Board of California.
27 Failure to pay such costs shall be considered a violation of probation.

28 Payment must be made in full within 30 calendar days of the effective date of the Order, or

1 by a payment plan approved by the Medical Board of California. Any and all requests for a
2 payment plan shall be submitted in writing by respondent to the Board. Failure to comply with
3 the payment plan shall be considered a violation of probation.

4 The filing of bankruptcy by respondent shall not relieve respondent of the responsibility to
5 repay investigation and enforcement costs, including expert review costs.

6 13. QUARTERLY DECLARATIONS. Respondent shall submit quarterly declarations
7 under penalty of perjury on forms provided by the Board, stating whether there has been
8 compliance with all the conditions of probation.

9 Respondent shall submit quarterly declarations not later than 10 calendar days after the end
10 of the preceding quarter.

11 14. GENERAL PROBATION REQUIREMENTS.

12 Compliance with Probation Unit

13 Respondent shall comply with the Board's probation unit.

14 Address Changes

15 Respondent shall, at all times, keep the Board informed of Respondent's business and
16 residence addresses, email address (if available), and telephone number. Changes of such
17 addresses shall be immediately communicated in writing to the Board or its designee. Under no
18 circumstances shall a post office box serve as an address of record, except as allowed by Business
19 and Professions Code section 2021, subdivision (b).

20 Place of Practice

21 Respondent shall not engage in the practice of medicine in Respondent's or patient's place
22 of residence, unless the patient resides in a skilled nursing facility or other similar licensed
23 facility.

24 License Renewal

25 Respondent shall maintain a current and renewed California physician's and surgeon's
26 license.

27 Travel or Residence Outside California

28 Respondent shall immediately inform the Board or its designee, in writing, of travel to any

1 areas outside the jurisdiction of California which lasts, or is contemplated to last, more than thirty
2 (30) calendar days.

3 In the event Respondent should leave the State of California to reside or to practice
4 Respondent shall notify the Board or its designee in writing 30 calendar days prior to the dates of
5 departure and return.

6 15. INTERVIEW WITH THE BOARD OR ITS DESIGNEE. Respondent shall be
7 available in person upon request for interviews either at Respondent's place of business or at the
8 probation unit office, with or without prior notice throughout the term of probation.

9 16. NON-PRACTICE WHILE ON PROBATION. Respondent shall notify the Board or
10 its designee in writing within 15 calendar days of any periods of non-practice lasting more than
11 30 calendar days and within 15 calendar days of Respondent's return to practice. Non-practice is
12 defined as any period of time Respondent is not practicing medicine as defined in Business and
13 Professions Code sections 2051 and 2052 for at least 40 hours in a calendar month in direct
14 patient care, clinical activity or teaching, or other activity as approved by the Board. If
15 Respondent resides in California and is considered to be in non-practice, Respondent shall
16 comply with all terms and conditions of probation. All time spent in an intensive training
17 program which has been approved by the Board or its designee shall not be considered non-
18 practice and does not relieve Respondent from complying with all the terms and conditions of
19 probation. Practicing medicine in another state of the United States or Federal jurisdiction while
20 on probation with the medical licensing authority of that state or jurisdiction shall not be
21 considered non-practice. A Board-ordered suspension of practice shall not be considered as a
22 period of non-practice.

23 In the event Respondent's period of non-practice while on probation exceeds 18 calendar
24 months, Respondent shall successfully complete the Federation of State Medical Board's Special
25 Purpose Examination, or, at the Board's discretion, a clinical competence assessment program
26 that meets the criteria of Condition 18 of the current version of the Board's "Manual of Model
27 Disciplinary Orders and Disciplinary Guidelines" prior to resuming the practice of medicine.

28 Respondent's period of non-practice while on probation shall not exceed two (2) years.

1 Periods of non-practice will not apply to the reduction of the probationary term.

2 Periods of non-practice for a Respondent residing outside of California will relieve
3 Respondent of the responsibility to comply with the probationary terms and conditions with the
4 exception of this condition and the following terms and conditions of probation: Obey All Laws;
5 General Probation Requirements; Quarterly Declarations; Abstain from the Use of Alcohol and/or
6 Controlled Substances; and Biological Fluid Testing.

7 17. COMPLETION OF PROBATION. Respondent shall comply with all financial
8 obligations (e.g., restitution, probation costs) not later than 120 calendar days prior to the
9 completion of probation. This term does not include cost recovery, which is due within 30
10 calendar days of the effective date of the Order, or by a payment plan approved by the Medical
11 Board and timely satisfied. Upon successful completion of probation, Respondent's certificate
12 shall be fully restored.

13 18. VIOLATION OF PROBATION. Failure to fully comply with any term or condition
14 of probation is a violation of probation. If Respondent violates probation in any respect, the
15 Board, after giving Respondent notice and the opportunity to be heard, may revoke probation and
16 carry out the disciplinary order that was stayed. If an Accusation, or Petition to Revoke Probation,
17 or an Interim Suspension Order is filed against Respondent during probation, the Board shall have
18 continuing jurisdiction until the matter is final, and the period of probation shall be extended until
19 the matter is final.

20 19. LICENSE SURRENDER. Following the effective date of this Decision, if
21 Respondent ceases practicing due to retirement or health reasons or is otherwise unable to satisfy
22 the terms and conditions of probation, Respondent may request to surrender his or her license.
23 The Board reserves the right to evaluate Respondent's request and to exercise its discretion in
24 determining whether or not to grant the request, or to take any other action deemed appropriate
25 and reasonable under the circumstances. Upon formal acceptance of the surrender, Respondent
26 shall within 15 calendar days deliver Respondent's wallet and wall certificate to the Board, or its
27 designee and Respondent shall no longer practice medicine. Respondent will no longer be subject
28 to the terms and conditions of probation. If Respondent re-applies for a medical license, the

1 application shall be treated as a petition for reinstatement of a revoked certificate.

2 20. PROBATION MONITORING COSTS. Respondent shall pay the costs associated
3 with probation monitoring each and every year of probation, as designated by the Board, which
4 may be adjusted on an annual basis. Such costs shall be payable to the Medical Board of
5 California and delivered to the Board or its designee no later than January 31 of each calendar
6 year.

7 21. FUTURE ADMISSIONS CLAUSE. If Respondent should ever apply or reapply for
8 a new license or certification, or petition for reinstatement of a license, by any other health care
9 licensing action agency in the State of California, all of the charges and allegations contained in
10 Accusation No. 800-2022-086193 shall be deemed to be true, correct, and admitted by
11 Respondent for the purpose of any Statement of Issues or any other proceeding seeking to deny or
12 restrict license.

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1 ACCEPTANCE

2 I have carefully read the above Stipulated Settlement and Disciplinary Order and have fully
3 discussed it with my attorney, Taryn A. Perez, Esq. I understand the stipulation and the effect it
4 will have on my Physician's and Surgeon's Certificate. I enter into this Stipulated Settlement and
5 Disciplinary Order voluntarily, knowingly, and intelligently, and agree to be bound by the
6 Decision and Order of the Medical Board of California.

7
8 DATED: 9/16/2025 PDT

F Omar B Tordilla, MD

9 Signature ID: JFEHCIXR76
F. OMAR BRIONES TORDILLA, M.D.
Respondent

10 I have read and fully discussed with Respondent F. Omar Briones Tordilla, M.D. the terms
11 and conditions and other matters contained in the above Stipulated Settlement and Disciplinary
12 Order. I approve its form and content.

13
14 DATED: 9/18/2025

Taryn A. Perez

15 TARYN A. PEREZ, ESQ.
Attorney for Respondent

16
17 ENDORSEMENT

18 The foregoing Stipulated Settlement and Disciplinary Order is hereby respectfully
19 submitted for consideration by the Medical Board of California.

20
21 DATED: 09/18/2025

Respectfully submitted,

22 ROB BONTA
Attorney General of California
23 MATTHEW M. DAVIS
Supervising Deputy Attorney General

24 Andres T. Carnahan

25 ANDRES T. CARNAHAN
26 Deputy Attorney General
27 Attorneys for Complainant
28

Exhibit A

Accusation No. 800-2022-086193

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7 Facsimile: (619) 645-2061

8 *Attorneys for Complainant*

9
10 **BEFORE THE**
MEDICAL BOARD OF CALIFORNIA
11 **DEPARTMENT OF CONSUMER AFFAIRS**
STATE OF CALIFORNIA

12
13 In the Matter of the Accusation Against:

Case No. 800-2022-086193

14 **F. OMAR BRIONES TORDILLA, M.D.**
225 E Second Ave Ste 101
15 Escondido, CA 92025

A C C U S A T I O N

16 **Physician's and Surgeon's Certificate**
No. A 88578,

17 Respondent.
18

19 **PARTIES**

20 1. Reji Varghese (Complainant) brings this Accusation solely in his official capacity as
21 the Executive Director of the Medical Board of California, Department of Consumer Affairs
22 (Board).

23 2. On or about August 13, 2004, the Medical Board issued Physician's and Surgeon's
24 Certificate Number A 88578 to F. Omar Briones Tordilla, M.D. (Respondent). The Physician's
25 and Surgeon's Certificate was in full force and effect at all times relevant to the charges brought
26 herein and will expire on November 30, 2025, unless renewed.

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4. Section 2004 of the Code states:

(a) The enforcement of the disciplinary and criminal provisions of the Medical Practice Act.

(c) Carrying out disciplinary actions appropriate to findings made by a panel or an administrative law judge.

(e) Reviewing the quality of medical practice carried out by physician and surgeon certificate holders under the jurisdiction of the board.

...

Except as otherwise provided by law, the board may take action against all persons guilty of violating this chapter. The board shall enforce and administer this article as to physician and surgeon certificate holders, including those who hold certificates that do not permit them to practice medicine, such as, but not limited to, retired, inactive, or disabled status certificate holders, and the board shall have all the powers granted in this chapter for these purposes . . .

(a) A licensee whose matter has been heard by an administrative law judge of the Medical Quality Hearing Panel as designated in Section 11371 of the Government Code, or whose default has been entered, and who is found guilty, or who has entered into a stipulation for disciplinary action with the board, may, in accordance with the provisions of this chapter:

(2) Have his or her right to practice suspended for a period not to exceed one year upon order of the board.

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1 (4) Be publicly reprimanded by the board. The public reprimand may include a
2 requirement that the licensee complete relevant educational courses approved by the
board.

3 (5) Have any other action taken in relation to discipline as part of an order of
4 probation, as the board or an administrative law judge may deem proper.

5 ...

6 STATUTORY PROVISIONS

7 7. Section 2234 of the Code states:

8 The board shall take action against any licensee who is charged with
9 unprofessional conduct. In addition to other provisions of this article, unprofessional
conduct includes, but is not limited to, the following:

10 (a) Violating or attempting to violate, directly or indirectly, assisting in or
11 abetting the violation of, or conspiring to violate any provision of this chapter.

12 (b) Gross negligence.

13 (c) Repeated negligent acts. To be repeated, there must be two or more
14 negligent acts or omissions. An initial negligent act or omission followed by a
separate and distinct departure from the applicable standard of care shall constitute
repeated negligent acts.

15 (1) An initial negligent diagnosis followed by an act or omission medically
16 appropriate for that negligent diagnosis of the patient shall constitute a single
negligent act.

17 (2) When the standard of care requires a change in the diagnosis, act, or
18 omission that constitutes the negligent act described in paragraph (1), including, but
19 not limited to, a reevaluation of the diagnosis or a change in treatment, and the
licensee's conduct departs from the applicable standard of care, each departure
constitutes a separate and distinct breach of the standard of care.

20 ...

21 8. Section 2266 of the Code states:

22 The failure of a physician and surgeon to maintain adequate and accurate
23 records relating to the provision of services to their patients constitutes unprofessional
conduct.

24 9. Section 2242 of the Code states:

25 (a) Prescribing, dispensing, or furnishing dangerous drugs as defined in Section
26 4022 without an appropriate prior examination and a medical indication, constitutes
unprofessional conduct. An appropriate prior examination does not require a
27 synchronous interaction between the patient and the licensee and can be achieved
through the use of telehealth, including, but not limited to, a self-screening tool or a
28 questionnaire, provided that the licensee complies with the appropriate standard of
care. ...

1 10. Section 4021 of the Code states:

2 “Controlled substance” means any substance listed in Chapter 2 (commencing
3 with Section 11053) of Division 10 of the Health and Safety Code.

4 11. Section 4022 of the Code states:

5 “Dangerous drug” ... means any drug ... unsafe for self-use in humans ... and
6 includes the following:

7 (a) Any drug that bears the legend: “Caution: federal law prohibits dispensing
8 without prescription,” “Rx only,” or words of similar import.

9 ...

10 (c) Any other drug ... that by federal or state law can be lawfully dispensed
11 only on prescription or furnished pursuant to Section 4006.

12 12. Unprofessional conduct under Business and Professions Code section 2234 is conduct
13 which breaches the rules or ethical code of the medical profession, or conduct which is
14 unbecoming to a member in good standing of the medical profession, and which demonstrates an
15 unfitness to practice medicine.¹

16 COST RECOVERY

17 13. Section 125.3 of the Code provides, in pertinent part, that the Board may request the
18 administrative law judge to direct a licensee found to have committed a violation or violations of
19 the licensing act to pay a sum not to exceed the reasonable costs of the investigation and
20 enforcement of the case, with failure of the licensee to comply subjecting the license to not being
21 renewed or reinstated. If a case settles, recovery of investigation and enforcement costs may be
22 included in a stipulated settlement.

23 DEFINITIONS

24 14. Percocet is a brand name for oxycodone and acetaminophen, a Schedule II controlled
25 substance pursuant to Health and Safety Code section 11055, subdivision (b), and a dangerous
26 drug pursuant to Business and Professions Code section 4022.

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¹ *Shea v. Board of Medical Examiners*, (1978) 81 Cal.App.3d 564, 575.

15. Norco is a brand name for a hydrocodone-acetaminophen combination product, a Schedule II controlled substance pursuant to Health and Safety Code section 11055, subdivision (b)(1)(I)(i), and a dangerous drug pursuant to Business and Professions Code section 4022.

16. Morphine milligram equivalents (MME) are values that represent the potency of an opioid dose relative to morphine. Using morphine as the standard, MME is a tool for doctors to compare different drugs in a simplified, unified measurement.

17. Ambien is a brand name for zolpidem tartrate, a Schedule IV controlled substance pursuant to Health and Safety Code section 11057, subdivision (d), and a dangerous drug pursuant to Business and Professions Code section 4022. It is a sedative used for the short-term treatment of insomnia.

18. Soma is a brand name for carisoprodol, a muscle relaxant. It is a Schedule IV controlled substance pursuant to Health and Safety Code section 11057, subdivision (d), and a dangerous drug pursuant to Code section 4022.

19. Flexeril is a brand name for cyclobenzaprine, a muscle relaxant. It is a dangerous drug pursuant to Code section 4022 but not a controlled substance.

STANDARD OF CARE

20. The standard of care for a provider in the state of California when transitioning a patient from acute opioid therapy to chronic opioid therapy (greater than 90 days) is to have a diagnosis of medical necessity.

21. The standard of care for a primary care provider in California, when considering long-term use of opioids for chronic non-cancer pain, given the potential risks of opioid analgesics, is to undertake risk stratification.

22. The standard of care for a primary care provider in the state of California, for patients who are continued on controlled substances after an initial trial, is to base care on outcomes such as making progress toward functional goals (activity level), presence and nature of side effects (adverse effects), pain status (analgesia), mental health status (affect), and any evidence of aberrant behavior such as patient misuse, abuse, or diversion. These are also often referred to as

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1 the five A's or five objectives in pain management: analgesia, activity level, adverse effects,
2 affect, and aberrant behaviors.

3 23. The standard of care for a primary care provider in California is, when considering
4 long-term use of opioids for chronic non-cancer pain, the provider and patient should develop
5 treatment goals. The goals of treatment should include: improvement in pain and function;
6 improvement in pain-associated symptoms such as sleep disturbance and anxiety/depression; and
7 additionally should include an exit strategy in the event it becomes medically necessary.

8 24. There is poor evidence for the use of opioids for muscle-skeletal pain and greater risk
9 for harm, especially when using chronic moderate-dose opioids when safer alternatives exist.

10 25. The standard of care when prescribing Soma is to use only for short periods, up to
11 three weeks.

12 26. The standard of care for a primary care provider in California, when considering the
13 long-term use of controlled substances, is that the physician discusses the risks and benefits of the
14 treatment plan with the patient.

15 27. The potential risks of long-term opioid use or chronic Soma use – and the combined
16 use of opioids and Soma – might include respiratory depression, motor impairment, cognitive
17 impairment, and death, and may lead to dependence, misuse, addiction and overdose.

18 28. The standard of care for the contents of medical records in California is that medical
19 records contain adequate documentation. Medical records serve as the basis for planning and
20 maintaining the quality of patient care. When devoid of important medical information, illegible
21 or unintelligible, the provider and other treating health professionals may remain unaware of
22 important aspects of a patient's medical condition.

23 29. The standard of care for a primary provider in the state of California, when
24 prescribing chronic opioids or other controlled substances, is to ensure appropriate compliance
25 monitoring.

26 30. Since October 2, 2018, providers have been required to consult CURES prior to
27 prescribing, ordering, administering, or furnishing a Schedule II-IV controlled substance. The

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1 law has since been amended to include Schedule V controlled substances. Providers must consult
2 CURES:

3 a. The first time² a patient is prescribed, ordered, administered, or furnished a controlled
4 substance, unless one of the exemptions apply.

5 b. Within the twenty-four-hour period, or the previous business day, before prescribing,
6 ordering, administering, or furnishing a controlled substance, unless one of the exemptions apply.

7 c. Before subsequently prescribing a controlled substance, if previously exempt. At
8 least once every six months if the controlled substance remains a part of the patient's treatment
9 plan.

10 FACTUAL ALLEGATIONS

11 31. At all relevant times, Respondent was a family practice physician in San Diego.

12 Patient 1:³

13 32. Patient 1 established primary care with Respondent on or about October 5, 2018, at
14 the age of 62 years.

15 33. On or about October 21, 2018, following aortic valve replacement surgery, the
16 provider concerned had prescribed Patient 1 a thirty-day supply of oxycodone 5 mg, to be taken
17 one to two tablets every four hours for pain, as needed.

18 34. On or about January 26, 2020, Patient 1 presented to the Urgent Care for lower back
19 pain at a level that was noted as "5/10." Patient 1 was given Toradol and Solu-Medrol (a
20 nonsteroidal anti-inflammatory and a steroid) by injection and advised to follow up with his
21 primary care physician ("PCP") or go to the Emergency Room ("ER") if needed.

22 35. On or about January 28, 2020, Patient 1 presented to Respondent for an urgent care
23 follow up for his back pain. The injections had helped but Patient 1 reported that his pain had
24 returned. Respondent prescribed 60 x Norco 325/5 mg tablets, a 30-day supply (representing
25 10 MME daily.)

26 ² "First time" is defined as the initial occurrence in which a health care practitioner intends
27 to prescribe, order, administer, or furnish a controlled substance to a patient and has not
previously prescribed a controlled substance to the patient.

28 ³ Patient names are known to all parties but not disclosed for privacy reasons.

1 36. Patient 1 returned the following day. Patient 1's chart indicates that his lower back
2 pain was continuing despite the medication. Respondent ordered an X-ray and doubled
3 Patient 1's Norco dosage (20 MME), and noted that an MRI would be ordered if there was no
4 improvement.

5 37. Patient 1's X-ray showed mild lumbar degenerative changes without significant disc
6 space narrowing.

7 38. On or about January 31, 2020, Patient 1 reported continued pain with no relief from
8 the increased Norco. On the same date, Respondent documented that he ordered an MRI without
9 contrast of Patient 1's lumbar spine.

10 39. On or about February 3, 2020, Patient 1 filled a prescription from Respondent for 30
11 x Percocet 325/5 mg tablets, for a 10-day supply (26 MME).

12 40. On or about February 11, 2020, a chart note by Respondent indicates that Patient 1
13 was unable to work and still had back pain; he would be placed on disability.⁴ The MRI had
14 showed edema and possible osteomyelitis, and an MRI with contrast was ordered.

15 41. Also on or about February 11, 2020, Patient 1 signed a Controlled Substance Use
16 Agreement.⁵

17 42. On or about February 12, 2020, Patient 1 filled a prescription from Respondent for 90
18 x Percocet 325/5 mg tablets, for a 30-day supply.

19 43. On or about February 18, 2020, Patient 1 presented for an office visit with reports of
20 right foot pain. Respondent ordered X-rays and labs.

21 44. The X-ray report of Patient 1's right foot showed no acute findings. Patient 1's lab
22 results dated February 18, 2020, were negative for markers of inflammatory arthritis such as gout,
23 connective tissue disorders, or rheumatoid arthritis.⁶

24
25 ⁴ Patient 1 was initially placed on two months' disability leave (starting January 29, 2020),
which was later extended by a further two months, on or about March 26, 2020.

26 ⁵ A handwritten annotation, "Update 10/6/22" with an unknown set of initials appears on
27 the first page of this agreement.

28 ⁶ There do not appear to be more general markers of inflammation completed on this date,
such as C-Reactive Protein (CRP) or Erythrocyte Sedimentation Rate (ESR).

1 45. On or about February 20, 2020, Respondent prescribed steroids for Patient 1, who had
2 reported continuing pain, now from his back, leg, and foot.

3 46. On or about February 21, 2020, the results of Patient 1's MRI with contrast (of his
4 lumbar spine) were positive for osteomyelitis.⁷

5 47. On or about March 6, 2020, Patient 1 presented for an office visit. Respondent
6 prescribed him Percocet 325/10 mg tablets, to be taken three times per day as needed (45 MME),
7 for his acute bilateral lower back pain.

8 48. On or about May 22, 2020, Patient 1 was still on the same dosage of Percocet 325/10
9 mg. Patient 1 presented for an office visit, wanting to return to work. Patient 1 reported feeling
10 good and that he had been cleared by infectious diseases and cardiology.

11 49. On or about May 26, 2020, Respondent again provided Patient 1 a refill for the
12 Percocet 325/10 mg, three times daily.

13 50. On or about June 3, 2020, Patient 1's Percocet 325/10 mg dosage was decreased to
14 twice daily (30 MME).

15 51. Patient 1 presented for an office visit on or about October 30, 2020. In Respondent's
16 chart note for this visit, he states Patient 1 was negative for muscle-skeletal complaints. In his
17 physical exam, Respondent describes Patient 1's muscle-skeletal and neurological systems as
18 normal. In his assessment and plan, Respondent will refill Patient 1's pain medications for his
19 low back pain and also for his inflammatory polyarthritis.

20 52. There is no indication in Patient 1's chart of how the diagnosis of inflammatory
21 polyarthritis was made.

22 53. Patient 1 again presented for continuing refills in March, August, and December
23 2021, and April and August 2022.

24 54. In his chart note for Patient 1's office visit dated August 2, 2022, Respondent again
25 states Patient 1 was negative for muscle-skeletal complaints and that his muscle-skeletal and

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28 ⁷ Osteomyelitis is inflammation or swelling that occurs in the bone.

1 neurological systems were normal. Respondent refilled Patient 1's pain medication for his
2 inflammatory polyarthritis. Respondent states:

3 "[Patient 1] aware of risk. [C]onsider tapering down or pain management. ..."

4 55. Also on August 2, 2022, Respondent's assessment and plan for Patient 1 includes
5 "acute pain of right shoulder," with an order for X-rays to be taken. It is unclear where this
6 diagnosis originates as there is no discussion of Patient 1's shoulder in the history of present
7 illness ("HPI"). The shoulder X-ray results show "no acute radiographic findings" and mild
8 osteoarthritis.

9 56. On or about August 31, 2022, Patient 1 asks for – and is given – a refill prescription
10 for cyclobenzaprine, a non-controlled substance. It is unclear from Patient 1's chart why this
11 medication is prescribed.⁸

12 57. On or about October 6, 2022, a one-year CURES report was pulled for Patient 1's
13 controlled substance prescriptions and can be found in his chart.⁹ There is no indication that
14 Respondent reviewed this report and there is no reference to it in any chart notes. This is the only
15 CURES report found in Patient 1's chart.

16 58. In a chart note dated October 13, 2022, Respondent states Patient 1 is negative for
17 muscle-skeletal complaints. His physical exam showed low paraspinal tenderness bilaterally.
18 Respondent documents that he and Patient 1 discussed, at length, the prolonged use of pain
19 medication and that Patient 1 had agreed to tapering down the medication. A plan was made to
20 decrease Percocet to one daily in November 2022.

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24 ⁸ In a note dated November 8, 2018, an employee documents a call from Patient 1, in
25 which he said he had "a stiff neck and was wondering if he could get some muscle relaxers." A
26 prescription (of an unidentified medication, presumably cyclobenzaprine) was apparently
27 approved and sent to the pharmacy. Cyclobenzaprine 10 mg is listed as one of Patient 1's active
28 medications on subsequent chart notes, reportedly stopped on September 25, 2019. The next (and
final, until August 31, 2022) reference to cyclobenzaprine is in a chart note dated January 28,
2020, when Respondent prescribed Flexeril (a brand name for cyclobenzaprine), along with
Norco and ibuprofen, for Patient 1's low back pain.

⁹ Pages 2 and 3 of the three-page report are in Patient 1's chart. Page 1 is absent.

1 59. Patient 1 was prescribed two Percocet 325/10 mg tablets daily (30 MME) for a total
2 of two years and five months, from June 2020 through November 2022. This was more than two
3 years following resolution of Patient 1's osteomyelitis.

4 60. On or about November 5, 2022, Patient 1's Percocet 325/10 mg dosage was
5 decreased to one per day (15 MME). Patient 1 remained on this medication and dosage for four
6 months, until March 2023.

7 61. On or about March 16, 2023, Patient 1 presented for an office visit. Per Respondent's
8 chart note of that date, Patient 1 was "trying to stop [P]ercocet. Trying to hold off[f] on pain
9 meds." Respondent did not prescribe any further opioids to Patient 1.

10 62. In an interview with the Medical Board, Respondent stated that he did not do urine
11 drug screens¹⁰ because Patient 1 did not give him any indication for them. There is no evidence
12 that Respondent utilized other compliance monitoring techniques, such as pill counting.

13 63. Respondent's records for Patient 1 do not show the use of any screening tools to
14 evaluate Patient 1's potential risks of misuse of a controlled substance.

15 64. There is no evidence, from Respondent's records for Patient 1, that Respondent
16 evaluated Patient 1's progress toward any treatment objectives. There is no indication of the use
17 of a 1 to 10 pain scale, or any ongoing assessment that includes descriptions of the anatomical
18 location of pain, quality of pain, timing of pain, palliation, and provocation of pain.

19 65. There is no evidence that Respondent consistently evaluated treatment goals such as
20 Patient 1's activity level (functional goals), adverse effects (side effects), aberrant behaviors
21 (signs of drug or alcohol use, unsanctioned dose escalation, and early refill requests), and
22 patient's affect (changes to mood, depression or anxiety).

23 66. Respondent's chart for Patient 1 does not adequately demonstrate a discussion with
24 Patient 1 regarding the potential risks of long-term opioid use, including the risk of respiratory
25 depression, motor impairment, cognitive impairment, and death. There was also no clearly
26 documented discussion of the risk of dependence, misuse, addiction, overdose, and death.

27
28 ¹⁰ Patient 1's chart contains lab results for one urinalysis conducted on or about November
13, 2018, along with a complete blood count ("CBC") and other labs.

1 67. Respondent's notes for Patient 1 almost uniformly do not include a detailed history,
2 and lack information on the assessment and planning necessary to allow Respondent and other
3 providers to determine if appropriate, adequate, and safe treatment was being provided to
4 Patient 1.

5 68. Respondent did not specify measurable goals and objectives to evaluate Patient 1's
6 treatment progress. His chart notes for Patient 1 fail to show discernible improvement in pain and
7 associated symptoms during the treatment period. Respondent also did not include an exit
8 strategy for discontinuing controlled substances therapy in the event that tapering or termination
9 of controlled substances therapy became necessary.

10 Patient 2:

11 69. According to Respondent, Patient 2 established primary care with him in 2010.¹¹

12 70. In January 2018, Patient 2 was a 56-year-old female whose chronic conditions
13 included: chronic pain disorder, narcotic dependence, post inflammatory pulmonary fibrosis,
14 insomnia, status post bariatric surgery, malignant lymphomas (unspecified site), paresthesia of
15 foot (change in sensation), and lumbosacral radiculopathy. On or about January 2, 2018,
16 Patient 2 filled prescriptions for 90 x Soma, 90 x Percocet 325/7.5 mg, and 30 x Ambien, in each
17 case a thirty-day supply.

18 71. On or about February 1, 2018, Patient 2 presented for an office visit for medications.
19 In Respondent's chart note for this visit, he notes no joint swelling and muscle weakness, and
20 documents normal musculoskeletal system and normal neurologic system.

21 72. From February 2018 through July 2019, Patient 2 continued to receive Percocet
22 325/7.5 mg for a daily MME of 33.8.

23 73. In a chart note for a visit on or about August 13, 2019, Respondent documented that
24 Patient 2 "needs increase in meds." There is no indication in the chart note of why Patient 2
25 needed a dose increase. The chart note stated, further, that the Percocet was to be increased to
26 four times daily "for the next 2 months," then return to three times daily in October 2019.

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28 ¹¹ Records provided to, and reviewed by, the Board cover care and treatment provided to
Patient 2 during the period January 1, 2018, through June 27, 2023.

1 74. Patient 2 remained on the increased dose of Percocet for three and a half years, until
2 May 2023, when it was dropped back to three times daily.

3 75. Patient 2 had an X-ray of her left hip on or about August 20, 2019. The radiologist's
4 impression was "normal left hip."

5 76. Patient 2 had X-rays of her left hip and her left shoulder on or about January 8, 2021.
6 The radiologist's impression was "normal left hip" and "normal left shoulder."

7 77. Respondent's requests for an MRI for Patient 2 were denied and, on or about
8 January 20, 2021, Respondent and Patient 2 agreed that she would try physical therapy for her left
9 hip. There is no indication in the chart of whether Patient 2 ever received physical therapy or, if
10 so, whether it had any effect on her reported condition.

11 78. Patient 2 filled prescriptions for ninety (90) Soma 350 mg tablets monthly (three per
12 day) from at least February 2019 through March 2022. According to Patient 2's CURES report,
13 in April 2022, Patient 2 started filling prescriptions from Respondent for one hundred twenty
14 (120) tablets per month, or four per day. The reason for the increased prescription is not
15 documented in Patient 2's chart.¹²

16 79. On or about July 6, 2022, X-rays were taken of Patient 2's right and left shoulders.
17 Both showed mild tendinitis and mild degenerative changes "without acute osseous abnormality."

18 80. On or about August 21, 2022, Respondent received a letter from a major retail
19 pharmacy about the dangers of prescribing a combination of Soma, Ambien, and Percocet to
20 Patient 2. A handwritten (undated) annotation on the margin of the (typed) letter states, "no more
21 Soma. Just Flexeril [cyclobenzaprine]."

22 81. On or about August 22, 2022, a one-year CURES report was pulled for Patient 2's
23 controlled substance prescriptions and can be found in her chart. There is no indication that
24

25 ¹² Patient 2 called the office on or about March 23, 2022, for a refill of her Soma
26 prescription, three per day. On or about the same date, she filled her usual ninety tablets (for
27 three per day). In the chart note for an office visit on or about April 21, 2022, Patient 2's
28 medications are reported as including four Soma tablets per day. Respondent's plan and
assessment is to "continue current treatment," and, starting April 21, 2022, Patient 2 fills
prescriptions for four Soma tablets per day. No other mention of the increase in Soma dosage can
be found in Patient 2's chart.

1 Respondent reviewed this report and there is no reference to it in any chart notes. This is the only
2 CURES report found in Patient 2's chart.

3 82. On or about September 19, 2022, Patient 2 called Respondent's office and asked for
4 refills for her Percocet and Ambien prescriptions, and asked for a prescription for Flexeril. It is
5 unclear why she asked for Flexeril or whether the "no more Soma" was communicated to her at
6 any time. The note, documenting the phone call, includes the word "Cures" without further
7 elucidation.

8 83. On or about December 22, 2022, Respondent documented in his chart note that
9 Patient 2 was aware of the risk of prolonged pain medication use and should taper down.
10 Mention was made of a "poss[ible] pain management referral."

11 84. Patient 2 remained on Percocet 325/7.5 mg four times daily, for an approximate daily
12 MME of 45, until May 2023.

13 85. On or about April 28, 2023, a certified letter was delivered to Patient 2, asking her to
14 authorize the release of her medical records to the Division of Investigation. Patient 2 did not
15 respond to the letter.

16 86. Respondent's chart for Patient 2 documents communications with her on or about
17 May 15 and May 16, 2023, regarding medication refills. In the note dated May 16, 2023,
18 Respondent documents:

19 "discussed at length chronic pain med use. suggested tapering down. pt. agrees. will
20 decrease from Percocet ... qid # 120 to Percocet ... tid #90, also consider pain
management."

21 87. Respondent did not do any urine drug screens on Patient 2.

22 88. Respondent prescribed to Patient 2 a combination of Ambien, Soma, and Percocet for
23 at least a period of five years. Following the August 2022 letter from the retail pharmacy
24 regarding the dangers of this combination, Respondent switched from Soma to Flexeril in
25 September 2022, and continued prescribing Percocet and Ambien until at least October 2023.

26 89. Respondent's records for Patient 2 do not show the use of any screening tools to
27 evaluate Patient 2's potential risks of misuse of a controlled substance.

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1 90. There is no evidence, from Respondent's records for Patient 2, that Respondent
2 evaluated Patient 2's progress toward any treatment objectives. There is no indication of the use
3 of a 1 to 10 pain scale, or any ongoing assessment that includes descriptions of the anatomical
4 location of pain, quality of pain, timing of pain, palliation, and provocation of pain.

5 91. There is no evidence that Respondent consistently evaluated treatment goals such as
6 Patient 2's activity level (functional goals), adverse effects (side effects), aberrant behaviors
7 (signs of drug or alcohol use, unsanctioned dose escalation, and early refill requests), and
8 patient's affect (changes to mood, depression or anxiety).

9 92. Respondent's chart for Patient 2 does not adequately demonstrate a discussion with
10 Patient 2 regarding the potential risks of long-term opioid use, including the risk of respiratory
11 depression, motor impairment, cognitive impairment, and death. There was also no clearly
12 documented discussion of the risk of dependence, misuse, addiction, overdose, and death.

13 93. Respondent's notes for Patient 2 almost uniformly do not include a detailed history,
14 and lack information on the assessment and planning necessary to allow Respondent and other
15 providers to determine if appropriate, adequate, and safe treatment was being provided to
16 Patient 2.

17 94. Respondent did not specify measurable goals and objectives to evaluate Patient 2's
18 treatment progress. His chart notes for Patient 2 fail to show discernible improvement in pain and
19 associated symptoms during the treatment period. Respondent also did not include an exit
20 strategy for discontinuing controlled substances therapy in the event that tapering or termination
21 of controlled substances therapy became necessary.

22 95. During the roughly five and a half years of records reviewed, Patient 2 made fourteen
23 (14) office visits to Respondent and had six (6) telehealth visits with Respondent. In
24 Respondent's notes for each of these twenty (20) visits, under "Review of Systems," he
25 documents that Patient 2 was negative for joint swelling and muscle weakness.

26 96. Respondent documents a normal musculoskeletal exam at each of nine (9) visits
27 between January 2018 and November 2019. No physical exam or demonstration of movement is
28 documented on any of the telehealth visits.

1 97. A single Narcotic Use Agreement was found in Patient 2's chart, dated October 3,
2 2017, and covering Patient 2's use of Percocet.

3 Patient 3:

4 98. Patient 3 is an adult female born in 1978.

5 99. Respondent's chart note for an office visit by Patient 3 on or about March 14, 2019¹³,
6 lists Patient 3's chronic conditions as: chondromalacia of the left knee, synovitis of the right knee,
7 gastroesophageal reflux disease without esophagitis, morbid obesity, narcotic dependence, right
8 adrenal mass, thyroid nodule, dysmenorrhea, chronic back pain, migraine, abnormal mammogram
9 of right breast, and hypothyroidism.

10 100. From (at least) January 2019 through September 2023, Respondent treated Patient 3's
11 muscle-skeletal pain (chronic low back pain and inflammatory polyarthritis) with Percocet 325/10
12 mg, to be taken three per day (for a daily MME of 45), decreasing on May 11, 2023, to two per
13 day for a daily MME of 30. Patient 3's final prescription for narcotics was filled by the pharmacy
14 on September 5, 2023, and sold on September 7, 2023.

15 101. In addition to the Percocet, Respondent prescribed Soma 350 mg tablets, three daily,
16 for Patient 3 from (at least) January 2019 through July 2022.¹⁴

17 102. In the records reviewed for Patient 3, the first reference to the risks of prolonged
18 opioid use is found in the chart note dated October 3, 2022. On the same date, Patient 3 signed a
19 "Patient Agreement for Prescription Opioids (Narcotics)" in which potential side effects of
20 opioids were described and initialed by Patient 3.¹⁵

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24 ¹³ The medical records reviewed for Patient 3 cover the period January 1, 2019, through
25 December 31, 2023.

26 ¹⁴ On or about August 21, 2022, Respondent received a letter from a major retail
pharmacy about the dangers of prescribing a combination of Soma, Ambien, and Percocet.

27 ¹⁵ Patient 3 signed a Controlled Substance Use Agreement on or about August 26, 2019, in
28 which her duties and responsibilities with regard to her controlled substance medications are
outlined.

1 103. Respondent did not do urinalysis on Patient 3. Patient 3's chart contains two CURES
2 reports, one obtained on or about October 3, 2022, and a second one on or about September 5,
3 2023.

4 104. Respondent's records for Patient 3 do not show the use of any screening tools to
5 evaluate Patient 3's potential risks of misuse of a controlled substance.

6 105. There is no evidence, from Respondent's records for Patient 3, that Respondent
7 evaluated Patient 3's progress toward any treatment objectives. There is no indication of the use
8 of a 1 to 10 pain scale, or any ongoing assessment that includes descriptions of the anatomical
9 location of pain, quality of pain, timing of pain, palliation, and provocation of pain.

10 106. There is no evidence that Respondent consistently evaluated treatment goals such as
11 Patient 3's activity level (functional goals), adverse effects (side effects), aberrant behaviors
12 (signs of drug or alcohol use, unsanctioned dose escalation, and early refill requests), and
13 patient's affect (changes to mood, depression or anxiety).

14 107. Respondent's chart for Patient 3 does not adequately demonstrate a discussion with
15 Patient 3 regarding the potential risks of long-term opioid use or chronic Soma use and/or the
16 combined use of opioids and Soma, including respiratory depression, motor impairment,
17 cognitive impairment, and death. There was also no clearly documented discussion of the risk of
18 dependence, misuse, addiction, overdose, and death.

19 108. Respondent's notes for Patient 3 almost uniformly do not include a detailed history,
20 and lack information on the assessment and planning necessary to allow Respondent and other
21 providers to determine if appropriate, adequate, and safe treatment was being provided to
22 Patient 3.

23 109. Respondent did not specify measurable goals and objectives to evaluate Patient 3's
24 treatment progress. His chart notes for Patient 3 fail to show discernible improvement in pain and
25 associated symptoms during the treatment period. Respondent also did not include an exit
26 strategy for discontinuing controlled substances therapy in the event that tapering or termination
27 of controlled substances therapy became necessary.

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1 Patient 4:

2 110. Patient 4 is an adult male, born in 1964. He is the husband of Patient 3.

3 111. Respondent's chart note for an office visit by Patient 4 on or about April 4, 2019¹⁶,
4 lists Patient 4's chronic conditions: morbid obesity, localized osteoarthritis of right knee,
5 inflammatory polyarthritis, arthritis and osteoarthritis of left knee, bilateral chronic knee pain,
6 narcotic dependence, sleep apnea, nocturnal hypoxia, nocturnal hypoxemia.

7 112. The chart note for the April 4, 2019, visit states, under "History of Present Illness,"
8 that Patient 4 "needs refills." No additional information is provided. All the findings from the
9 physical exam of Patient 4 are documented as normal, and Patient 4's muscle-skeletal system
10 shows no joint swelling or muscle weakness.

11 113. On or about February 11, 2019, Patient 4 saw orthopedic surgeon, Dr. B., in a follow-
12 up appointment for his left knee. According to the chart note for that visit, Patient 4 reported his
13 pain level as 3 on a scale of 1 to 10.

14 114. On or about July 20, 2022, Patient 4 saw Dr. B. for a consult of his right hip.
15 According to the chart note for that visit, Patient 4 reported his pain level as 1 to 2 out of 10.
16 Patient 4 reported that his symptoms had decreased significantly over the prior couple of weeks.
17 In comparing X-rays, Dr. B. found minimal progression in the arthritic changes in Patient 4's hip
18 over a period of four years.

19 115. Like Patient 3, from (at least) January 2019 through September 2023, Respondent
20 treated Patient 4's muscle-skeletal pain (bilateral knee pain and inflammatory polyarthritis) with
21 Percocet 325/10 mg. Patient 4's dosage was approximately 60 MME until May 11, 2023, when it
22 was decreased to 45 MME. Patient 4's final prescription for narcotics was filled by the pharmacy
23 on September 5, 2023, and sold on September 7, 2023.

24 116. In addition, like with Patient 3, Respondent prescribed Soma 350 mg tablets, three
25 daily, to Patient 4, from (at least) January 2019 through June 2022.

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28 ¹⁶ The medical records reviewed for Patient 4 cover the period January 1, 2019, through
December 31, 2023.

1 117. With rare exceptions, for the entirety of the period reviewed, Patient 4's Percocet and
2 Soma prescriptions were filled and/or sold on the same date as Patient 3's Percocet and Soma
3 prescriptions.

4 118. Respondent did not do urinalysis on Patient 4.¹⁷

5 119. Patient 4's chart contains four CURES reports, two of which were obtained on or
6 about the same date as Patient 3's CURES report(s), namely, October 3, 2022, and September 5,
7 2023. In addition, there are patient CURES reports for Patient 4 dated June 24, 2022, and August
8 26, 2023.

9 120. Like Patient 3, Patient 4 signed a Controlled Substance Use Agreement on or about
10 August 26, 2019.

11 121. Like Patient 3, Patient 4 signed a Patient Agreement for Prescription Opioids
12 (Narcotics) on or about October 3, 2022.

13 122. Respondent's records for Patient 4 do not show the use of any screening tools to
14 evaluate Patient 4's potential risks of misuse of a controlled substance.

15 123. There is no evidence, from Respondent's records for Patient 4, that Respondent
16 evaluated Patient 4's progress toward any treatment objectives. There is no indication of the use
17 of a 1 to 10 pain scale, or any ongoing assessment that includes descriptions of the anatomical
18 location of pain, quality of pain, timing of pain, palliation, and provocation of pain.

19 124. There is no evidence that Respondent consistently evaluated treatment goals such as
20 Patient 4's activity level (functional goals), adverse effects (side effects), aberrant behaviors
21 (signs of drug or alcohol use, unsanctioned dose escalation, and early refill requests), and
22 patient's affect (changes to mood, depression or anxiety).

23 125. Respondent's chart for Patient 4 does not adequately demonstrate a discussion with
24 Patient 4 regarding the potential risks of long-term opioid use or chronic Soma use and/or the
25 combined use of opioids and Soma, including respiratory depression, motor impairment,

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27 ¹⁷ According to a note made following a phone call with Patient 4 on or about September
28 5, 2023 (the date of Patient 4's final prescription for narcotics), a urine drug screen was ordered.
There is no indication in the chart of whether it was performed and, if so, its results.

1 cognitive impairment, and death. There was also no clearly documented discussion of the risk of
2 dependence, misuse, addiction, overdose, and death.

3 126. Respondent's notes for Patient 4 almost uniformly do not include a detailed history,
4 and lack information on the assessment and planning necessary to allow Respondent and other
5 providers to determine if appropriate, adequate, and safe treatment was being provided to
6 Patient 4.

7 127. Respondent did not specify measurable goals and objectives to evaluate Patient 4's
8 treatment progress. His chart notes for Patient 4 fail to show discernible improvement in pain and
9 associated symptoms during the treatment period. Respondent also did not include an exit
10 strategy for discontinuing controlled substances therapy in the event that tapering or termination
11 of controlled substances therapy became necessary.

12 Patient 5:

13 128. Patient 5, a male adult, was born in 1955. He was a patient of Respondent, who
14 prescribed Patient 5 opioids from at least 2007 through February 2023.¹⁸

15 129. Respondent treated Patient 5's muscle-skeletal pain (chronic low back pain,
16 inflammatory polyarthritis, and "chronic pain") with a combination of Norco 325/5 mg and
17 Percocet 325/10 mg¹⁹ continuously from at least January 2018 through August 2022, for a
18 continuous, combined MME of approximately 90 to 102.

19 130. Respondent discontinued the Percocet in August 2022, but continued prescribing
20 Norco 325/5 (13 MME) to Patient 5 until February 2023.

21 131. According to a patient CURES report for Patient 5, he filled his prescriptions at
22 multiple pharmacies and paid by various methods, including commercial insurance, private pay,
23 and "other." Patient 5 requested early refills of his opioids on multiple occasions.

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25 ¹⁸ Medical records were reviewed for Patient 5 covering the period January 2018 through
26 March 2023.

27 ¹⁹ Respondent also prescribed Percocet 325/10 mg, four tablets per day (60 MME) to
28 Patient 5's wife, continuously from at least January 2019 through April 2023, and to Patient 5's
son continuously from at least January 2019 through July 2019, when he passed away due to
acute fentanyl and oxycodone intoxication.

1 132. On or about October 5, 2018, a representative from CVS pharmacy called
2 Respondent's office to confirm that Patient 5 was to receive both Percocet and Norco. According
3 to the telephone message documented in Patient 5's chart, the CVS employee stated that since
4 Patient 5, Patient 5's son and Patient 5's spouse were all getting opioid pain medications, it might
5 be best for them to be referred to pain management instead of getting their medications from
6 Respondent, their primary care provider.

7 133. On or about January 11, 2019, a representative from CVS pharmacy again expressed
8 concern about dispensing both Percocet and Norco to Patient 5.

9 134. On or about February 11, 2019, Patient 5 reported further problems with filling
10 prescriptions for both Percocet and Norco at CVS. Reportedly, the pharmacist insisted that
11 Patient 5 would benefit from pain management and refused to refill more than a seven-day supply
12 of Percocet to hold him over. Patient 5 requested that his future prescriptions be sent to
13 Walgreens pharmacy.

14 135. On or about February 25, 2019, Patient 5 asked for his Norco prescriptions to be sent
15 to Indian Health Pharmacy.

16 136. On or about March 1, 2019, a representative of Indian Health Pharmacy contacted
17 Respondent's office for confirmation that Patient 5 was going to be on both Norco and Percocet
18 simultaneously.

19 137. A patient CURES report for Patient 5 was pulled on April 30, 2019, and marked
20 "Reviewed."

21 138. On or about October 2, 2019, Patient 5 signed a Controlled Substance Use
22 Agreement.

23 139. On or about June 5, 2020, a representative from Walgreens pharmacy called
24 Respondent's office to confirm that Patient 5 was receiving both Percocet and Norco medications.

25 140. On or about December 30, 2021, Patient 5 called Respondent's office and asked that
26 his Norco refill be sent to Rite Aid pharmacy as he is no longer using Walgreens. On the same
27 day, a representative from Rite Aid Pharmacy called Respondent's office saying they did not feel
28 comfortable filling the Norco prescription since Patient 5 also had a prescription for Percocet.

1 141. Per a note in Patient 5's chart dated December 30, 2021, Patient 5 told Respondent's
2 office staff that the Rite Aid pharmacist had declined to fill his medication because he did not
3 have pharmacy benefits. He asked that the Norco prescription be sent to Indian Health Pharmacy.

4 142. On or about July 14, 2022, Patient 5 called Respondent's office and asked for his
5 Percocet prescription to be transferred to Sav-On pharmacy. He called twice again on the same
6 date, to follow up on the transfer request. On or about July 15, 2022, Patient 5 called again about
7 transferring the refill order of his Percocet to Sav-On, and roughly an hour and a half later,
8 Patient 5's son called about the same transfer request.

9 143. On or about July 21, 2022, Patient 5 told Respondent's office staff that the pharmacist
10 at Sav-On would only give him a one-week's supply of Percocet at a time, until Patient 5 saw a
11 pain management physician.

12 144. Patient 5 had a pain management appointment on August 11, 2022. On or about
13 August 8, 2022, Patient 5 called and said it was imperative that he speak with Respondent before
14 August 11, 2022. On or about August 9, 2022, Patient 5 again called to say that he was going to
15 cancel his pain management appointment and would stop taking Percocet. He asked for a refill of
16 his Norco prescription to be sent to Indian Health.

17 145. A note in Patient 5's chart dated September 12, 2022, states that Respondent spoke to
18 Patient 5 "at length concerning his norco." Patient 5 had been sent to pain management but did
19 "not want to see them." Patient 5 only wanted to see Respondent.

20 "[Patient 5] already has stopped his percocet. [Patient 5] aware of risk with
21 cont[ained] norco use. discussed the amount of norco he is taking #180. [Patient 5] is
22 will[ing] to taper down and decrease amount of norco 5/325 #120. [Patient 5] is on
contract. refill his norco 5/325 #120."

23 146. A patient CURES report for Patient 5 was pulled on September 12, 2022, and
24 initialed.²⁰

25 147. According to Patient 5's chart, Respondent discussed with him the risks associated
26 with prolonged pain medication use on or about November 16, 2022. In a note dated February

27 ²⁰ Patient 5's chart for the period reviewed contained two CURES reports, pulled April 30,
28 2019, and September 12, 2022. Besides these two reports, a further report dated August 27, 2015
(outside the period of review) was found in Patient 5's chart.

1 22, 2023, Respondent noted that Patient 5 was "aware of prolonged pain med[ication] use."
2 Patient 5 "refused to go back [to pain management]. Consider tapering down."

3 **FIRST CAUSE FOR DISCIPLINE**

4 **(Gross Negligence)**

5 148. Respondent is subject to disciplinary action under sections 2227 and 2234, as defined
6 by section 2234, subdivision (b), of the Code, in that he committed gross negligence in his care
7 and treatment of Patient 1, Patient 2, Patient 3, Patient 4, and Patient 5, as more particularly
8 alleged hereinafter:

9 149. Paragraphs 14 through 147, above, are hereby realleged and incorporated by this
10 reference as if fully set forth herein.

11 150. Patient 1:

12 a) Between approximately May 2020 through at least November 2022, Respondent used
13 chronic opioid therapy on Patient 1 in the absence of a diagnosis of medical necessity and/or to
14 treat muscle-skeletal pain.

15 b) Respondent failed to classify Patient 1's risk prior to initiating his long-term use of
16 opioids.

17 c) Respondent failed to utilize any objectives to fully evaluate Patient 1's controlled
18 substances needs.

19 d) Respondent failed to elucidate a coherent treatment plan and objectives when
20 utilizing chronic opioid therapy on Patient 1.

21 e) Respondent failed to clearly elucidate to Patient 1 the long-term risks or side effects
22 of opioids.

23 f) Respondent failed to conduct adequate compliance monitoring to ensure Patient 1's
24 appropriate and/or safe use of controlled substances.

25 g) Respondent failed to keep adequate medical records of his care and treatment of
26 Patient 1.

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151. Patient 2:

- a) Respondent treated Patient 2 with chronic moderate-dose opioids without an appropriate medical indication.
- b) Respondent prescribed to Patient 2 for chronic use Soma, Percocet, and Ambien – three medications that, when used concurrently, are synergistic for negative health outcomes.
- c) Respondent failed to utilize the five objectives in pain management to fully evaluate Patient 2's controlled substance needs and/or determine the efficacy of the treatment.
- d) Respondent prescribed to Patient 2 a sedative/hypnotic not indicated for long-term therapy, namely, Ambien, for a period of at least six and a half years.
- e) Respondent failed to utilize treatment guidelines when prescribing Soma to Patient 2, when safer alternatives were available.
- f) Respondent failed to elucidate clear treatment goals and objectives in his ongoing prescribing of controlled substances to Patient 2.
- g) Respondent failed to clearly elucidate to Patient 2 the risks or side effects of opioids, sedative/hypnotics, and Soma, and their combined use.
- h) Respondent failed to adequately undertake advised compliance monitoring of Patient 2 to meet patient safety goals and/or legal requirements.
- i) Respondent failed to keep adequate medical records on his care and treatment of Patient 2.

152. Patient 3:

- a) Respondent treated Patient 3 with chronic moderate-dose opioids without an appropriate medical indication.
- b) Respondent prescribed to Patient 3 for chronic use Soma and Percocet – two medications that, when used concurrently, are synergistic for negative health outcomes.
- c) Respondent failed to utilize treatment guidelines when prescribing Soma to Patient 3, when safer alternatives were available.
- d) Respondent failed to develop any treatment plan or objectives when utilizing chronic opioid therapy in his care and treatment of Patient 3.

1 e) Respondent failed to utilize the five objectives in pain management to evaluate
2 Patient 3's controlled substance needs and/or determine the efficacy of the treatment.

3 f) Respondent failed to clearly elucidate to Patient 3 the risks or side effects of opioids,
4 sedative/hypnotics, and Soma, and their combined use.

5 g) Respondent failed to adequately undertake advised compliance monitoring of
6 Patient 3 to meet patient safety goals and/or legal requirements.

7 h) Respondent failed to keep adequate medical records on his care and treatment of
8 Patient 3.

9 153. Patient 4:

10 a) Respondent treated Patient 4 with chronic moderate-dose opioids without an
11 appropriate medical indication.

12 b) Respondent prescribed to Patient 4 for chronic use Soma and Percocet - two
13 medications that, when used concurrently, are synergistic for negative health outcomes.

14 c) Respondent failed to utilize treatment guidelines when prescribing Soma to Patient 4,
15 when safer alternatives were available.

16 d) Respondent failed to develop any treatment plan or objectives when utilizing chronic
17 opioid therapy in his care and treatment of Patient 4.

18 e) Respondent failed to utilize the five objectives in pain management to evaluate
19 Patient 4's controlled substance needs and/or determine the efficacy of the treatment.

20 f) Respondent failed to clearly elucidate to Patient 4 the risks or side effects of opioids,
21 sedative/hypnotics, and Soma, and their combined use.

22 g) Respondent failed to adequately undertake advised compliance monitoring of
23 Patient 4 to meet patient safety goals and/or legal requirements.

24 h) Respondent failed to keep adequate medical records on his care and treatment of
25 Patient 4.

26 154. Patient 5:

27 a) Respondent treated Patient 5 with chronic high-dose opioids without an appropriate
28 medical indication.

1 b) Respondent failed to develop any treatment plan or objectives when utilizing chronic
2 opioid therapy in his care and treatment of Patient 5.

3 c) Respondent failed to utilize the five objectives in pain management to evaluate
4 Patient 5's controlled substance needs and/or determine the efficacy of the treatment.

5 d) Respondent failed to clearly and timely elucidate to Patient 5 the risks or side effects
6 of opioids, sedative/hypnotics, and their combined use.

7 e) Respondent failed to adequately undertake advised compliance monitoring of
8 Patient 5 to meet patient safety goals and/or legal requirements.

9 f) Respondent failed to keep adequate medical records on his care and treatment of
10 Patient 5.

11 **SECOND CAUSE FOR DISCIPLINE**

12 **(Repeated Negligent Acts)**

13 155. Respondent is further subject to disciplinary action under sections 2227 and 2234, as
14 defined by section 2234, subdivision (c), of the Code, in that he committed repeated negligent
15 acts in his care and treatment of Patient 1, Patient 2, Patient 3, Patient 4, and Patient 5, as more
16 particularly alleged in paragraphs 14 through 154, above, which are hereby realleged and
17 incorporated by this reference as if fully set forth herein.

18 **THIRD CAUSE FOR DISCIPLINE**

19 **(Prescribing Dangerous Drugs Without Prior Examination and Medical Indication)**

20 156. Respondent is further subject to disciplinary action under sections 2227 and 2234, as
21 defined by section 2242, of the Code, in that he prescribed dangerous drugs as defined in Section
22 4022 of the Code without an appropriate prior examination and a medical indication, as more
23 particularly alleged in paragraphs 14 through 155, above, which are hereby realleged and
24 incorporated by reference as if fully set forth herein.

25 **FOURTH CAUSE FOR DISCIPLINE**

26 **(Failure to Maintain Adequate and Accurate Records)**

27 157. Respondent is further subject to disciplinary action under sections 2227 and 2234, as
28 defined by section 2266, of the Code, in that he failed to maintain adequate and accurate records

1 relating to his provision of services to Patient 1, Patient 2, Patient 3, Patient 4, and Patient 5, as
2 more particularly alleged in paragraphs 14 through 155, above, which are hereby realleged and
3 incorporated by this reference as if fully set forth herein.

4 **FIFTH CAUSE FOR DISCIPLINE**

5 **(General Unprofessional Conduct)**

6 158. Respondent is further subject to disciplinary action under 2234 of the Code, in that he
7 has engaged in conduct which breaches the rules or ethical code of the medical profession, or
8 conduct that is unbecoming to a member in good standing of the medical profession, and which
9 demonstrates an unfitness to practice medicine, as more particularly alleged in paragraphs 14
10 through 157, above, which are hereby realleged and incorporated by this reference as if fully set
11 forth herein.

12 **PRAYER**

13 WHEREFORE, Complainant requests that a hearing be held on the matters herein alleged,
14 and that following the hearing, the Medical Board of California issue a decision:

- 15 1. Revoking or suspending Physician's and Surgeon's Certificate Number A 88578,
16 issued to Respondent F. Omar Briones Tordilla, M.D.;
- 17 2. Revoking, suspending or denying approval of Respondent F. Omar Briones Tordilla,
18 M.D.'s authority to supervise physician assistants and advanced practice nurses;
- 19 3. Ordering Respondent F. Omar Briones Tordilla, M.D., to pay the Board the costs of
20 the investigation and enforcement of this case, and if placed on probation, the costs of probation
21 monitoring; and
- 22 4. Taking such other and further action as deemed necessary and proper.

23 DATED: JAN 08 2025

24 Jenna Jones For
25 REJI VARGHESE
26 Executive Director
27 Medical Board of California
28 Department of Consumer Affairs
State of California
Complainant