

LBD00303-05-6

1 1. "CONSUMER" MEANS THE PERSON UPON WHICH A MEDICAL DEVICE WAS USED,
2 ATTACHED OR APPLIED, REGARDLESS OF WHO PURCHASED OR ACQUIRED SUCH
3 DEVICE.

4 2. "ELECTRONIC MEDICAL DEVICE" MEANS AN IMPLANTABLE MEDICAL DEVICE
5 THAT REQUIRES A BATTERY OR SIMILAR POWER SOURCE TO FUNCTION.

6 3. "IMPLANTABLE HIP OR KNEE MEDICAL DEVICE" MEANS A DEVICE THAT
7 REPLACES THE COMPONENTS OF A HIP OR KNEE.

8 4. "INITIAL SELLER" MEANS THE SELLER WHO MANUFACTURED, MODIFIED,
9 REBUILT, IMPROVED OR RECONDITIONED AN IMPLANTABLE ELECTRONIC OR HIP OR
10 KNEE MEDICAL DEVICE.

11 S 646. EXPRESS WARRANTY REQUIRED. 1. EVERY INITIAL SELLER OF AN ELEC-
12 TRONIC MEDICAL DEVICE OR IMPLANTABLE HIP OR KNEE MEDICAL DEVICE SHALL
13 PROVIDE EACH CONSUMER OF SUCH DEVICE WITH A WARRANTY THAT THE MEDICAL
14 DEVICE IS FIT FOR THE ORDINARY PURPOSES FOR WHICH SUCH DEVICE IS USED,
15 AND IS FREE FROM DEFECTS FOR A PERIOD OF AT LEAST FIVE YEARS AFTER THE
16 MEDICAL DEVICE IS FIRST USED BY, ATTACHED TO OR APPLIED TO THE CONSUMER.

17 2. IF A MEDICAL DEVICE FAILS TO CONFORM TO THE WARRANTY REQUIRED BY
18 SUBDIVISION ONE OF THIS SECTION, AND THE CONSUMER, OR HIS OR HER AUTHOR-
19 IZED REPRESENTATIVE REPORTS SUCH NONCONFORMITY OR DEFECT TO THE INITIAL
20 SELLER OR ITS AGENTS DURING THE TERM OF THE WARRANTY, THE INITIAL SELLER
21 SHALL BE LIABLE FOR ALL COSTS INCURRED BY THE CONSUMER OR HIS OR HER
22 INSURER TO MAKE SUCH REPAIRS AND REPLACEMENTS AS ARE NECESSARY TO
23 CORRECT SUCH CONFORMITY OR DEFECT, AND ANY ADDITIONAL MEDICAL AND REHA-
24 BILITATION CARE NECESSARY AFTER SUCH REPAIR OR REPLACEMENT.

25 S 647. ADDITIONAL REMEDIES OF CONSUMERS. NOTHING IN THIS ARTICLE SHALL
26 IN ANY WAY LIMIT THE RIGHTS, REMEDIES OR PRIVILEGES WHICH ARE OTHERWISE
27 AVAILABLE TO A CONSUMER AT LAW OR EQUITY.

28 S 648. PROHIBITION AGAINST WAIVER OF RIGHTS. WAIVER OF ANY RIGHTS BY
29 THE CONSUMER UNDER THIS ARTICLE SHALL BE DEEMED CONTRARY TO PUBLIC POLI-
30 CY AND SHALL BE UNENFORCEABLE AND VOID.

31 S 649. EXCLUSION. THE PROVISIONS OF THIS ARTICLE SHALL NOT APPLY TO
32 DEVICES APPROVED THROUGH THE UNITED STATES FOOD AND DRUG ADMINISTRATION
33 PRE-MARKET APPROVAL PROCESS WHERE 21 USC 360(K) WOULD PROHIBIT IMPOSI-
34 TION OF THE WARRANTY ESTABLISHED UNDER THIS ARTICLE.

35 S 3. This act shall take effect on the first of January next succeed-
36 ing the date on which it shall have become a law, and shall apply to
37 medical devices initially used by, attached to or applied to a person on
38 or after such date.