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From: Tim Harrington  
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Re: FCC ID: YSGKA960  
Applicant: SORIN CRM  
Correspondence Reference Number: 39722  
Form 731 Confirmation Number: EA237591  
Date of Original E-mail: 04/22/2011

TO: Agent

CC: Applicant

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Please address the following items to facilitate continued FCC application processing.

1) a) as we understand this application is for a programmer / controller / monitor transmitting device intended to communicate with a 95 subpart I implant transmitter

- we note that attachments mention "IMD" i.e. implant transmitter

- fyi note that complete separate form-731 and 2.1033(c) and maybe 2.1033(b) contents AND a separate different FCC ID are required for an implant transmitter device; i.e. this application does not pertain to equipment auth. requirements for any implant transmitter devices

[This application is not an implant transmitter device. This monitor communicates with an IMD.](#)  
[This monitor communicates with an IMD.](#)

[This equipment object of this application is a monitor transmitting device intended to communicate with a 95 subpart I implant transmitter. This is not an implant transmitter device.](#)

[MedRadio is use in all documentation in place of "IMD"](#)

1) b) although this application was submitted as form-731 eqpt. class TNT "transmitter worn on body", based on info at hand it does not appear this device has intended nor reasonably expected worn-on-body and/or body-adjacent or held-on-body transmitting configurations - or if it does, please provide details about such use configurations if such body use is not applicable, fyi we will change eqpt. class to be TNB (or PCB, depending on response to item 2 herein)

[OK I think that the class is TNB and form-731 has been modified.](#)

1) c) fyi note that eqpt. auth. for 95 subpart I devices such as this that are not implant transmitters can be processed by TCBs [(<http://www.fcc.gov/oet/ea/procedures.html#sec3>), ([http://www.fcc.gov/oet/ea/ea\\_app\\_info.html](http://www.fcc.gov/oet/ea/ea_app_info.html))]; in contrast implant transmitters (with associated numerical electromagnetics modelling data to demonstrate compliance with RF exposure SAR limits) must submit application at FCC not TCB

[I propose to continue this certification with FCC](#)

1) d) if you want to continue with processing of this application as filed at FCC rather than a TCB, the other items in this corresp. need to be addressed accordingly; alternatively in reply you could request that FCC dismiss this application, after which the fee paid to FCC would be refunded and you could contact a TCB to arrange processing and under their independent fee structure

We want to continue with processing of this application as filed at FCC rather than a TCB,

2) as submitted this form-731 has only one rule-part/freq. line entry, i.e. for 95 subpart I; however other contents indicate device contains also 15.247 functions, which has its own specific equipment auth. and filing requirements especially, and related to 47 CFR 2.1033(e), for devices operating under both licensed-service [2.1033(c)] and part 15 [2.1033(b)] rules, two or more form-731 applications are required, in accordance with the following considerations

Yes, device contains also 15.247 functions, Ok I note that it has its own specific equipment auth. and filing requirements especially, other form-731 applications will be submitted.

3) as we understand the 15.247 function/component is an integral part of this device, and device will always be marketed and operated with the 15.247 component installed for the 15.247 portion, an f-731 under eqpt. class DTS or DSS must be submitted, depending on which sections of 15.247 are applicable in that case this FCC ID YSGKA961 represents the combination of at minimum the 95-I and 15.247 components / functions

Yes, 15.247 is applicable, I suggest that FCC ID YSGKA960 represents the combination of at minimum the 95-I and 15.247 components / functions. 2 applications form-731 have been completed.

4) please explain what are MedRadio and 15.247 transmit paths and components in the block diag. exhibit (i.e. f-731 exhibit type 4, presently has 1-page .pdf), and/or revise where appropriate to show those

There are included in the same equipment and PCB. The block Diagram has been modified.

(<https://apps.fcc.gov/oetcf/eas/misc/EasFaq.cfm>)

Question: What are the exhibit types required for an electronic filing?

Answer: There are 13 different exhibit types; they are listed below: (The number of exhibits required for a submittal is dependent upon the type of equipment for which authorization is sought.)

1. ID Label/Location Info
2. Attestation Statements
3. External Photos
4. Block Diagrams
5. Schematics
6. Test Report
7. Test Setup Photos
8. User's Manual
9. Internal Photos

10. Parts List/Tune-up Info

11. RF Exposure Info

12. Operational Description

13. Cover Letters

These 13 items have been sent (see last mail on November 4<sup>th</sup> 2011)

5) concerning confidentiality request cover letter in this filing, fyi note that in general internal photos exhibits do not qualify for permanent confidentiality, as described in e.g. KDB pub. 726290; please revise accordingly

Done

6) Some attachments mention "MICS"

- fyi since approx. May 2009 MICS is renamed to be MedRadio; therefore "MICS" is generally not applicable

Done

7) per 15.101(b), the receiver function in a MedRadio transceiver is subject to Verification eqpt. auth., which then requires also inclusion of the 15.19(a)(1) statement labelled on the device or in the documentation [15.19(a)(5)] - please explain compliance and/or revise exhibit(s) where appropriate

OK, done in the documentation.

fyi note this requirement is separate from the 15.105(b) statements required for Class B digital devices

Ok

8) emc/radio report pg 4 mentions "OKK for 2400-2483.5" - please explain what is OKK, or revise where appropriate

OKK was a mistake, modification done in test report.

9) for pg 14 of emc/radio report, because limit uses unit of microwatt, please revise to include listing of results as converted to microwatt

Ok

The items indicated above must be submitted before processing can continue on the above referenced application. Failure to provide the requested information within 60 days of the original e-mail date may result in application dismissal and forfeiture of the filing fee pursuant to Section 1.1108

DO NOT Reply to this email by using the Reply button. In order for your response to be processed expeditiously, you must upload your response via the Internet at [www.fcc.gov](http://www.fcc.gov), E-Filing, OET Equipment Authorization Electronic Filing, Submit Correspondence, Select Correspondence pertaining to EAS (Equipment Authorization System). Also, please note that partial responses increase processing time and should not be submitted.

Any questions about the content of this correspondence should be directed to the email address listed below the name of the sender.