LOGIQ V5/LOGIQ V3

Basic Service Manual



Part Number: 5496012-100 Revision: 3

Important Precautions

WARNING (EN)	 THIS SERVICE MANUAL IS AVAILABLE IN ENGLISH ONLY. IF A CUSTOMER'S SERVICE PROVIDER REQUIRES A LANGUAGE OTHER THAN ENGLISH, IT IS THE CUSTOMER'S RESPONSIBILITY TO PROVIDE TRANSLATION SERVICES. DO NOT ATTEMPT TO SERVICE THE EQUIPMENT UNLESS THIS SERVICE MANUAL HAS BEEN CONSULTED AND IS UNDERSTOOD. FAILURE TO HEED THIS WARNING MAY RESULT IN INJURY TO THE SERVICE PROVIDER, OPERATOR OR PATIENT FROM ELECTRIC SHOCK, MECHANICAL OR OTHER HAZARDS.
AVERTISSEMENT (FR)	 CE MANUEL DE MAINTENANCE N'EST DISPONIBLE QU'EN ANGLAIS. SI LE TECHNICIEN DU CLIENT A BESOIN DE CE MANUEL DANS UNE AUTRE LANGUE QUE L'ANGLAIS, C'EST AU CLIENT QU'IL INCOMBE DE LE FAIRE TRADUIRE. NE PAS TENTER D'INTERVENTION SUR LES ÉQUIPEMENTS TANT QUE LE MANUEL SERVICE N'A PAS ÉTÉ CONSULTÉ ET COMPRIS. LE NON-RESPECT DE CET AVERTISSEMENT PEUT ENTRAÎNER CHEZ LE TECHNICIEN, L'OPÉRATEUR OU LE PATIENT DES BLESSURES DUES à DES DANGERS ÉLECTRIQUES, MÉCANIQUES OU AUTRES.
WARNUNG (DE)	 DIESES KUNDENDIENST-HANDBUCH EXISTIERT NUR IN ENGLISCHER SPRACHE. FALLS EIN FREMDER KUNDENDIENST EINE ANDERE SPRACHE BENÖTIGT, IST ES AUFGABE DES KUNDEN FÜR EINE ENTSPRECHENDE ÜBERSETZUNG ZU SORGEN. VERSUCHEN SIE NICHT, DAS GERÄT ZU REPARIEREN, BEVOR DIESES KUNDENDIENST-HANDBUCH NICHT ZU RATE GEZOGEN UND VERSTANDEN WURDE. WIRD DIESE WARNUNG NICHT BEACHTET, SO KANN ES ZU VERLETZUNGEN DES KUNDENDIENSTTECHNIKERS, DES BEDIENERS ODER DES PATIENTEN DURCH ELEKTRISCHE SCHLÄGE, MECHANISCHE ODER SONSTIGE GEFAHREN KOMMEN.

ESTE MANUAL DE SERVICIO SÓ LO EXISTE EN INGLÉS.

• SI ALGÚN PROVEEDOR DE SERVICIOS AJENO A GEHC SOLICITA UN IDIOMA QUE NO SEA EL INGLÉS, ES RESPONSABILIDAD DEL CLIENTE OFRECER UN SERVICIO DE TRADUCCIÓN.



- NO SE DEBERÁ DAR SERVICIO TÉCNICO AL EQUIPO, SIN HABER CONSULTADO Y COMPRENDIDO ESTE MANUAL DE SERVICIO.
- LA NO OBSERVANCIA DEL PRESENTE AVISO PUEDE DAR LUGAR A QUE EL PROVEEDOR DE SERVICIOS, EL OPERADOR O EL PACIENTE SUFRAN LESIONES PROVOCADAS POR CAUSAS ELÉCTRICAS, MECÁNICAS O DE OTRA NATURALEZA.

ESTE MANUAL DE ASSISTÊNCIA TÉCNICA SÓ SE ENCONTRA DISPONÍVEL EM INGLÊS.

- SE QUALQUER OUTRO SERVIÇO DE ASSISTÊNCIA TÉCNICA, QUE NÃO A GEHC, SOLICITAR ESTES MANUAIS NOUTRO IDIOMA, É DA RESPONSABILIDADE DO CLIENTE EORNECER OS SERVIDOS DE TRADUÇÃO
- RESPONSABILIDADE DO CLIENTE FORNECER OS SERVIÇOS DE TRADUÇÃO.
- NãO TENTE REPARAR O EQUIPAMENTO SEM TER CONSULTADO E COMPREENDIDO ESTE MANUAL DE ASSISTÊNCIA TÉCNICA.
- O NãO CUMPRIMENTO DESTE AVISO PODE POR EM PERIGO A SEGURANÇA DO TÉCNICO, OPERADOR OU PACIENTE DEVIDO A' CHOQUES ELÉTRICOS, MECÂNICOS OU OUTROS.

ESTE MANUAL DE ASSISTÊNCIA ESTÁ DISPONÍVEL APENAS EM INGLÊS.

- SE QUALQUER OUTRO SERVIÇO DE ASSISTÊNCIA TÉCNICA, QUE NÃO A GEHC, SOLICITAR ESTES MANUAIS NOUTRO IDIOMA, É DA RESPONSABILIDADE DO CLIENTE FORNECER OS SERVIÇOS DE TRADUÇÃO.
- NÃO TENTE EFECTUAR REPARAÇÕES NO EQUIPAMENTO SEM TER CONSULTADO E COMPREENDIDO PREVIAMENTE ESTE MANUAL.
- A INOBSERVÂNCIA DESTE AVISO PODE RESULTAR EM FERIMENTOS NO TÉCNICO DE ASSISTÊNCIA, OPERADOR OU PACIENTE EM CONSEQUÊNCIA DE CHOQUE ELÉCTRICO, PERIGOS DE ORIGEM MECÂNICA, BEM COMO DE OUTROS TIPOS.

IL PRESENTE MANUALE DI MANUTENZIONE È DISPONIBILE SOLTANTO IN INGLESE.

- SE UN ADDETTO ALLA MANUTENZIONE ESTERNO ALLA GEHC RICHIEDE IL MANUALE IN UNA LINGUA DIVERSA, IL CLIENTE È TENUTO A PROVVEDERE DIRETTAMENTE ALLA TRADUZIONE.
- SI PROCEDA ALLA MANUTENZIONE DELL'APPARECCHIATURA SOLO DOPO AVER CONSULTATO IL PRESENTE MANUALE ED AVERNE COMPRESO IL CONTENUTO.
 - NON TENERE CONTO DELLA PRESENTE AVVERTENZA POTREBBE FAR COMPIERE OPERAZIONI DA CUI DERIVINO LESIONI ALL'ADDETTO ALLA MANUTENZIONE, ALL'UTILIZZATORE ED AL PAZIENTE PER FOLGORAZIONE ELETTRICA, PER URTI MECCANICI OD ALTRI RISCHI.

-

ATENÇÃO (PT-Br)



AVVERTENZA

(IT)

KÄESOLEV TEENINDUSJUHEND ON SAADAVAL AINULT INGLISE KEELES. KUI KLIENDITEENINDUSE OSUTAJA Nõ UAB JUHENDIT INGLISE KEELEST ERINEVAS KEELES. VASTUTAB KLIENT TÕLKETEENUSE OSUTAMISE EEST. HOIATUS **äRGE ÜRITAGE SEADMEID TEENINDADA ENNE EELNEVALT KÄESOLEVA** (ET) TEENINDUSJUHENDIGA TUTVUMIST JA SELLEST ARU SAAMIST. Kä ESOLEVA HOIATUSE EIRAMINE VÕIB PÕHJUSTADA TEENUSEOSUTAJA. OPERAATORI VÕI PATSIENDI VIGASTAMIST ELEKTRILÖÖGI. MEHAANILISE VõI MUU OHU TAGAJä RJEL. TÄMÄ HUOLTO-OHJE ON SAATAVILLA VAIN ENGLANNIKSI. • JOS ASIAKKAAN PALVELUNTARJOAJA VAATII MUUTA KUIN ENGLANNINKIELISTÄ MATERIAALIA, TARVITTAVAN KÄÄNNÖKSEN HANKKIMINEN ON ASIAKKAAN VASTUULLA. VAROITUS ä Lä YRITä KORJATA LAITTEISTOA ENNEN KUIN OLET VARMASTI LUKENUT (FI) JA YMMÄRTÄNYT TÄMÄN HUOLTO-OHJEEN. • MIKä LI TÄ TÄ VAROITUSTA EI NOUDATETA, SEURAUKSENA VOI OLLA PALVELUNTARJOAJAN. LAITTEISTON KÄYTTÄ JÄN TAI POTILAAN VAHINGOITTUMINEN SÄHKÖISKUN, MEKAANISEN VIAN TAI MUUN VAARATILANTEEN VUOKSI. ΤΟ ΠΑΡΟΝ ΕΓΧΕΙΡΙΔΙΟ ΣΕΡΒΙΣ ΔΙΑΤΙΘΕΤΑΙ ΣΤΑ ΑΓΓΛΙΚΑ ΜΟΝΟ. ΕΑΝ ΤΟ ΑΤΟΜΟ ΠΑΡΟΧΗΣ ΣΕΡΒΙΣ ΕΝΟΣ ΠΕΛΑΤΗ ΑΠΑΙΤΕΙ ΤΟ ΠΑΡΟΝ ΕΓΧΕΙΡΙΔΙΟ ΣΕ ΓΛΩΣΣΑ ΕΚΤΟΣ ΤΩΝ ΑΓΓΛΙΚΩΝ, ΑΠΟΤΕΛΕΙ ΕΥΘΥΝΗ ΤΟΥ ΠΕΛΑΤΗ ΝΑ ΠΑΡΕΧΕΙ ΥΠΗΡΕΣΙΕΣ ΜΕΤΑΦΡΑΣΗΣ. ΜΗΝ ΕΠΙΧΕΙΡΗΣΕΤΕ ΤΗΝ ΕΚΤΕΛΕΣΗ ΕΡΓΑΣΙΩΝ ΣΕΡΒΙΣ ΣΤΟΝ ΕΞΟΠΛΙΣΜΟ ΠΡΟΕΙΔΟΠΟΙΗΣΗ ΕΚΤΟΣ ΕΑΝ ΕΧΕΤΕ ΣΥΜΒΟΥΛΕΥΤΕΙ ΚΑΙ ΕΧΕΤΕ ΚΑΤΑΝΟΗΣΕΙ ΤΟ ΠΑΡΟΝ (EL) ΕΓΧΕΙΡΙΔΙΟ ΣΕΡΒΙΣ. ΕΑΝ ΔΕ ΛΑΒΕΤΕ ΥΠΟΨΗ ΤΗΝ ΠΡΟΕΙΔΟΠΟΙΗΣΗ ΑΥΤΗ, ΕΝΔΕΧΕΤΑΙ ΝΑ ΠΡΟΚΛΗΘΕΙ ΤΡΑΥΜΑΤΙΣΜΟΣ ΣΤΟ ΑΤΟΜΟ ΠΑΡΟΧΗΣ ΣΕΡΒΙΣ. ΣΤΟ ΧΕΙΡΙΣΤΗ Ή ΣΤΟΝ ΑΣΘΕΝΗ ΑΠΟ ΗΛΕΚΤΡΟΠΛΗΞΙΑ, ΜΗΧΑΝΙΚΟΥΣ Ή ΑΛΛΟΥΣ ΚΙΝΔΥΝΟΥΣ. EZEN KARBANTARTÁSI KÉZIKÖNYV KIZÁRÓLAG ANGOL NYELVEN ÉRHETŐ EL. HA A VEVŐ SZOLGÁLTATÓJA ANGOLTÓL ELTÉRŐ NYELVRE TART IGÉNYT. AKKOR A VEVŐ FELELŐSSÉGE A FORDÍTÁS ELKÉSZÍTTETÉSE. NE PRÓBÁLJA ELKEZDENI HASZNÁLNI A BERENDEZÉST, AMÍG A FIGYELMEZTETÉS KARBANTARTÁSI KÉZIKÖNYVBEN LEÍRTAKAT NEM ÉRTELMEZTÉK. (HU) EZEN FIGYELMEZTETÉS FIGYELMEN KÍVÜL HAGYÁSA A SZOLGÁLTATÓ, MŰKÖDTETŐ VAGY A BETEG ÁRAMÜTÉS. MECHANIKAI VAGY EGYÉB

VESZÉLYHELYZET MIATTI SÉRÜLÉSÉT EREDMÉNYEZHETI.

ÞESSI ÞJÓNUSTUHANDBÓK ER EINGÖNGU FÁANLEG Á ENSKU. EF ÞJÓNUSTUAÐILI VIÐSKIPTAMANNS ÞARFNAST ANNARS TUNGUMÁLS EN ENSKU. ER ÞAÐ Á ÁBYRGÐ VIÐSKIPTAMANNS AÐ ÚTVEGA ÞÝÐINGU. REYNIÐ EKKI AÐ ÞJÓNUSTA TÆKIÐ NEMA EFTIR AÐ HAFA SKOÐAÐ OG VIÐVÖRUN SKILIÐ ÞESSA ÞJÓNUSTUHANDBÓK. (IS) EF EKKI ER FARIÐ AÐ ÞESSARI VIÐVÖRUN GETUR ÞAÐ VALDIÐ MEIÐSLUM ÞJÓNUSTUVEITANDA. STJÓRNANDA EÐA SJÚKLINGS VEGNA RAFLOSTS. VÉLRÆNNAR EÐA ANNARRAR HÆTTU. TENTO SERVISNÍ NÁVOD EXISTUJE POUZE V ANGLICKÉM JAZYCE. • V Př (PADě, ŽE POSKYTOVATEL SLUŽEB ZÁKAZNÍKŮM POTř EBUJE Ná VOD V JINÉM JAZYCE. JE ZAJIŠTĚNÍ PŘ EKLADU DO ODPOVÍDAJÍCÍHO JAZYKA ú KOLEM Zá KAZNÍKA. NEPROVÁDĚJTE ú DRŽBU TOHOTO ZAŤ (ZENÍ, ANIŽ BYSTE SI PŤ EČ ETLI VÝSTRAHA TENTO SERVISNÍ NÁVOD A POCHOPILI JEHO OBSAH. (CS) V Př (PADě NEDODRŽOVANÍ TÉTO VýSTRAHY MůŽE DOJÍT ÚRAZU ELEKTRICKÁM PROUDEM PRACOVNÍKA POSKYTOVATELE SLUŽEB. OBSLUŽNÉHO PERSONÁLU NEBO PACIENTŮ VLIVEM ELEKTRICKÉHOP PROUDU, RESPEKTIVE VLIVEM K RIZIKU MECHANICKÉHO POŠKOZENÍ NEBO JINÉMU RIZIKU. DENNE SERVICEMANUAL FINDES KUN PÅ ENGELSK. HVIS EN KUNDES TEKNIKER HAR BRUG FOR ET ANDET SPROG END ENGELSK, ER DET KUNDENS ANSVAR AT SØRGE FOR OVERSÆTTELSE. FORSØG IKKE AT SERVICERE UDSTYRET MEDMINDRE ADVARSEL (DA) DENNE SERVICEMANUAL ER BLEVET LÆST OG FORSTÅET. MANGLENDE OVERHOLDELSE AF DENNE ADVARSEL KAN MEDFRE SKADE PL GRUND AF ELEKTRISK, MEKANISK ELLER ANDEN FARE FOR TEKNIKEREN, OPERATŘREN ELLER PATIENTEN. DEZE ONDERHOUDSHANDLEIDING IS ENKEL IN HET ENGELS VERKRIJGBAAR. ALS HET ONDERHOUDSPERSONEEL EEN ANDERE TAAL VEREIST. DAN IS DE KLANT VERANTWOORDELIJK VOOR DE VERTALING ERVAN. PROBEER DE APPARATUUR NIET TE ONDERHOUDEN VOORDAT DEZE WAARSCHUWING ONDERHOUDSHANDLEIDING WERD GERAADPLEEGD EN BEGREPEN IS. (NL) INDIEN DEZE WAARSCHUWING NIET WORDT OPGEVOLGD. ZOU HET ONDERHOUDSPERSONEEL. DE OPERATOR OF EEN PATIËNT GEWOND KUNNEN RAKEN ALS GEVOLG VAN EEN ELEKTRISCHE SCHOK. **MECHANISCHE OF ANDERE GEVAREN.**

ŠĪ APKALPES ROKASGRĀMATA IR PIEEJAMA TIKAI ANGĻU VALODĀ. JA KLIENTA APKALPES SNIEDZĒJAM NEPIECIEŠAMA INFORMĀCIJA CITĀ VALODĀ, NEVIS ANGĻU, KLIENTA PIENĀKUMS IR NODROŠINĀT TULKOŠANU. NEVEICIET APRĪKOJUMA APKALPI BEZ APKALPES ROKASGRĀMATAS BRĪDINĀJUMS IZLASĪŠANAS UN SAPRAŠANAS. (LV) ŠĪ BRĪDINĀJUMA NEIEVĒROŠANA VAR RADĪT ELEKTRISKĀS STRĀVAS TRIECIENA. MEHĀNISKU VAI CITU RISKU IZRAISĪTU TRAUMU APKALPES SNIEDZĒJAM, OPERATORAM VAI PACIENTAM. ŠIS EKSPLOATAVIMO VADOVAS YRA IŠLEISTAS TIK ANGLŲ KALBA. JEI KLIENTO PASLAUGU TEIKĖJUI REIKIA VADOVO KITA KALBA – NE ANGLU. VERTIMU PASIRŪPINTI TURI KLIENTAS. • NEMĖGINKITE ATLIKTI ĮRANGOS TECHNINĖS PRIEŽIŪROS DARBŲ, NEBENT ISPĖJIMAS VADOVAUTUMĖTĖS ŠIUO EKSPLOATAVIMO VADOVU IR JI SUPRASTUMĖTE (LT) NEPAISANT ŠIO PERSPĖJIMO, PASLAUGŲ TEIKĖJAS, OPERATORIUS AR PACIENTAS GALI BŪTI SUŽEISTAS DĖL ELEKTROS SMŪGIO. MECHANINIU AR **KITU PAVOJU.** DENNE SERVICEHÅNDBOKEN FINNES BARE PÅ ENGELSK. HVIS KUNDENS SERVICELEVERANDØR TRENGER ET ANNET SPRÅK, ER DET KUNDENS ANSVAR Å SØRGE FOR OVERSETTELSE. IKKE FORSØK Å REPARERE UTSTYRET UTEN AT DENNE. ADVARSEL (NO) SERVICEHÅNDBOKEN ER LEST OG FORSTÅTT. MANGLENDE HENSYN TIL DENNE ADVARSELEN KAN FØRE TIL AT SERVICELEVERANDØREN. OPERATØREN ELLER PASIENTEN SKADES PÅ GRUNN AV ELEKTRISK STØT, MEKANISKE ELLER ANDRE FARER. NINIEJSZY PODRĘCZNIK SERWISOWY DOSTĘPNY JEST JEDYNIE W JĘZYKU ANGIELSKIM. JEŚ LI FIRMA ŚWIADCZĄCA KLIENTOWI USŁUGI SERWISOWE WYMAGA UDOSTę PNIENIA PODRę CZNIKA W Ję ZYKU INNYM NIŻ ANGIELSKI, OBOWIAZEK ZAPEWNIENIA STOSOWNEGO TłUMACZENIA SPOCZYWA NA KLIENCIE. OSTRZEŻENIE NIE PRÓ BOWAĆ SERWISOWAĆ NINIEJSZEGO SPRZĘTU BEZ UPRZEDNIEGO (PL) ZAPOZNANIA SIĘ Z PODRĘCZNIKIEM SERWISOWYM. NIEZASTOSOWANIE SIę DO TEGO OSTRZEŻENIA MOŻE GROZIĆ OBRAŻENIAMI CIAłA SERWISANTA, OPERATORA LUB PACJENTA W WYNIKU PORAŻENIA PRĄDEM, URAZU MECHANICZNEGO LUB INNEGO RODZAJU ZAGROŻEń.

ACEST MANUAL DE SERVICE ESTE DISPONIBIL NUMAI ÎN LIMBA ENGLEZĂ. DACĂ UN FURNIZOR DE SERVICII PENTRU CLIENTI NECESITĂ O ALTĂ LIMBĂ DECÂT CEA ENGLEZĂ, ESTE DE DATORIA CLIENTULUI SĂ FURNIZEZE O TRADUCERE. ATENŢIE NU ÎNCERCATI SĂ REPARATI ECHIPAMENTUL DECÂT ULTERIOR (RO) CONSULTĂRII ȘI ÎNȚELEGERII ACESTUI MANUAL DE SERVICE. IGNORAREA ACESTUI AVERTISMENT AR PUTEA DUCE LA RĂNIREA DEPANATORULUI, OPERATORULUI SAU PACIENTULUI ÎN URMA PERICOLELOR DE ELECTROCUTARE, MECANICE SAU DE ALTĂ NATURĂ. Данное руководство по обслуживанию ПРЕДОСТАВЛЯЕТСЯ только на английском Языке. • Если сервисно МУ ПЕРСОНАЛУ клиента необходимо руководство не на английском ЯЗЫКЕ, клиенту следует самостоЯтельно ОБЕСПЕЧИТЬ перевод. ОСТОРОЖНО! • ПЕРЕД ОБСЛУЖИВАНИЕМ ОБОРУДОВАНИЯ ОБЯЗАТЕЛЬНО ОБРАТИТЕСЬ (RU) К ДАННОМУ РУКОВОДСТВУ И ПОЙМИТЕ ИЗЛОЖЕННЫЕ В НЕМ СВЕДЕНИЯ. НЕСОБЛЮДЕНИЕ УКАЗАННЫХ ТРЕБОВАНИЙ МОЖЕТ ПРИВЕСТИ К ТОМУ. ЧТО СПЕЦИАЛИСТ ПО ТЕХОБСЛУЖИВАНИЮ. ОПЕРАТОР ИЛИ ПАЦИЕНТ ПОЛУЧАТ УДАР ЗЛЕКТРИЧЕСКИМ ТОКОМ, МЕХАНИЧЕСКУЮ ТРАВМУ ИЛИ ДРУГОЕ ПОВРЕЖДЕНИЕ. ТОВА СЕРВИЗНО РЪКОВОДСТВО Е НАЛИЧНО САМО НА АНГЛИЙСКИ ЕЗИК. АКО ДОСТАВЧИКЪТ НА СЕРВИЗНИ УСЛУГИ НА КЛИЕНТ СЕ НУЖДАЕ ОТ ЕЗИК, РАЗЛИЧЕН ОТ АНГЛИЙСКИ, ЗАДЪЛЖЕНИЕ НА КЛИЕНТА Е ДА ПРЕДОСТАВИ ПРЕВОДАЧЕСКА УСЛУГА. НЕ СЕ ОПИТВАЙТЕ ДА ИЗВЪРШВАТЕ СЕРВИЗНО ОБСЛУЖВАНЕ НА ТОВА ПРЕДУПРЕЖДЕНИЕ ОБОРУДВАНЕ, ОСВЕН ВСЛУЧАЙ, ЧЕ СЕРВИЗНОТО РЪКОВОДСТВО Е ПРОЧЕТЕНО И СЕ РАЗБИРА. (BG) • НЕСПАЗВАНЕТО НА ТОВА ПРЕДУПРЕЖДЕНИЕ МОЖЕ ДА ДОВЕДЕ ДО НАРАНЯВАНЕ НА ДОСТАВЧИКА НА СЕРВИЗНИ УСЛУГИ, НА ОПЕРАТОРА ИЛИ ПАЦИЕНТА ВСЛЕДСТВИЕНА ТОКОВ УДАР, МЕХАНИЧНИ ИЛИ ДРУГИ РИСКОВЕ. OVAJ PRIRUČNIK ZA SERVISIRANJE DOSTUPAN JE SAMO NA ENGLESKOM JEZIKU. AKO KLIJENTOV SERVISER ZAHTEVA JEZIK KOJI NIJE ENGLESKI, ODGOVORNOST JE NA KLIJENTU DA PRUŽI USLUGE PREVOĐENJA. UPOZORENJE NEMOJTE POKUŠAVATI DA SERVISIRATE OPREMU AKO NISTE PROČITALI I (SR) RAZUMELI PRIRUČNIK ZA SERVISIRANJE. AKO NE POŠTUJETE OVO UPOZORENJE, MOŽE DOĆI DO POVREĐIVANJA SERVISERA, OPERATERA ILI PACIJENTA UZROKOVANOG ELEKTRIČNIM UDAROM, MEHANIČKIM I DRUGIM OPASNOSTIMA.

OPOZORILO (SL)	 TA SERVISNI PRIROČNIK JE NA VOLJO SAMO V ANGLEŠČINI. ČE PONUDNIK SERVISNIH STORITEV ZA STRANKO POTREBUJE NAVODILA V DRUGEM JEZIKU, JE ZA PREVOD ODGOVORNA STRANKA SAMA. NE POSKUŠAJTE SERVISIRATI OPREME, NE DA BI PREJ PREBRALI IN RAZUMELI SERVISNI PRIROČNIK. ČE TEGA OPOZORILA NE UPOŠTEVATE, OBSTAJA NEVARNOST ELEKTRIČNEGA UDARA, MEHANSKIH ALI DRUGIH NEVARNOSTI IN POSLEDIČNIH POŠKODB PONUDNIKA SERVISNIH STORITEV, UPORABNIKA OPREME ALI PACIENTA.
UPOZORENJE (HR)	 OVAJ SERVISNI PRIRUČNIK DOSTUPAN JE SAMO NA ENGLESKOM JEZIKU. AKO KLIJENTOV SERVISER ZAHTIJEVA JEZIK KOJI NIJE ENGLESKI, ODGOVORNOST KLIJENTA JE PRUŽITI USLUGE PREVOĐENJA. NEMOJTE POKUŠAVATI SERVISIRATI OPREMU AKO NISTE PROČITALI I RAZUMJELI SERVISNI PRIRUČNIK. AKO NE POŠTUJETE OVO UPOZORENJE, MOŽE DOĆI DO OZLJEDE SERVISERA, OPERATERA ILI PACIJENTA PROUZROČENE STRUJNIM UDAROM, MEHANIČKIM I DRUGIM OPASNOSTIMA.
UPOZORNENIE (SK)	 TÁTO SERVISNÁ PRÍRUČKA JE K DISPOZÍCII LEN V ANGLIČTINE. AK ZÁKAZNÍKOV POSKYTOVATEĽ SLUŽIEB VYŽADUJE INÝ JAZYK AKO ANGLIČTINU, POSKYTNUTIE PREKLADATEĽSKÝCH SLUŽIEB JE ZODPOVEDNOSŤOU ZÁKAZNÍKA. NEPOKÚŠAJTE SA VYKONÁVAŤ SERVIS ZARIADENIA SKÔR, AKO SI NEPREČÍTATE SERVISNÚ PRÍRUČKU A NEPOROZUMIETE JEJ. ZANEDBANIE TOHTO UPOZORNENIA Mô ŽE VYÚSTIŤ DO ZRANENIA POSKYTOVATEĽA SLUŽIEB, OBSLUHUJÚ CEJ OSOBY ALEBO PACIENTA ELEKTRICKÝM PRÚDOM, PRÍPADNE DO MECHANICKÉHO ALEBO INÉHO NEBEZPEČ ENSTVA.
VARNING (SV)	 DEN HÄR SERVICEHANDBOKEN FINNS BARA TILLGÄNGLIG PÅ ENGELSKA. OM EN KUNDS SERVICETEKNIKER HAR BEHOV AV ETT ANNAT SPRÅK ÄN ENGELSKA ANSVARAR KUNDEN FÖR ATT TILLHANDAHÅLLA ÖVERSÄTTNINGSTJÄNSTER. FÖRSÖK INTE UTFÖRA SERVICE PÅ UTRUSTNINGEN OM DU INTE HAR LÄST OCH FÖRSTÅR DEN HÄR SERVICEHANDBOKEN. OM DU INTE TAR HÄNSYN TILL DEN HÄR VARNINGEN KAN DET RESULTERA I SKADOR PÅ SERVICETEKNIKERN, OPERATÖREN ELLER PATIENTEN TILL FÖLJD AV ELEKTRISKA STÖTAR, MEKANISKA FAROR ELLER ANDRA FAROR.

BU SERVİS KILAVUZU YALNIZCA İNGİLİZCE OLARAK SAĞLANMIŞTIR.

 EĞER MÜŞTERİ TEKNİSYENİ KILAVUZUN İNGİLİZCE DIŞINDAKİ BİR DİLDE OLMASINI İSTERSE, KILAVUZU TERCÜME ETTİRMEK MÜŞTERİNİN SORUMLULUĞUNDADIR.

DİKKAT (TR)

- SERVİS KILAVUZUNU OKUYUP ANLAMADAN EKİPMANLARA MÜDAHALE ETMEYİNİZ.
- BU UYARININ GÖZ ARDI EDİLMESİ, ELEKTRİK ÇARPMASI YA DA MEKANİK VEYA DİĞER TÜRDEN KAZALAR SONUCUNDA TEKNİSYENİN, OPERATÖRÜN YA DA HASTANIN YARALANMASINA YOL AÇABİLİR.

このサービスマニュアルには英語版しかありません。

GEHC 以外でサービスを担当される業者が英語以外の言語を要求される場合、翻訳作業はその業者の責任で行うものとさせていただきます。

このサービスマニュアルを熟読し理解せずに、装置のサービスを行わないで下さい。

この警告に従わない場合、サービスを担当される方、操作員あるいは 患者さんが、感電や機械的又はその他の危険により負傷する可能性が あります。

本服務手冊僅提供英文版。

- 如顧客之服務提供者需要英文版以外之語言, 顧客需自行負擔其 翻譯服務之責任。
- •在查閱並了解本服務手冊之內容前,請勿試圖維修本設備。
- 未確實遵守本警告,可能導致服務提供者、操作者或病患遭受電撃、 機械危險或其他傷害。



本维修手册仅存有英文本・

非 GEHC 公司的维修员要求非英文本的维修手册时, 客户需自行负责翻译。

未详细阅读和完全了解本手册之前,不得进行维修。 忽略本注意事项会对维修员,操作员或病人造成触 电,机械伤害或其他伤害。



/干亰

(ZH-CN)

- 본 서비스 지침서는 영어로만 이용하실 수 있습니다.
- ·고객의 서비스 제공자가 영어이외 언어를 요구할 경우, 번역 서비스 지침서를 제공하는 것은 고객의 책임입니다.
- · 본 서비스 지침서를 지참했고 이해하지 않는 한은 해당 장비를 수리를 시도하지 마십시오.

-

·이 경우에 유해하지 않은 전기쇼크, 기계상의 혹은 다른 위험으로부터 서비스 제공자, 운영자 혹은 환자에게 위험을 가할 수 있습니다.

DAMAGE IN TRANSPORTATION

All packages should be closely examined at time of delivery. If damage is apparent write "Damage In Shipment" on ALL copies of the freight or express bill BEFORE delivery is accepted or "signed for" by a GE representative or hospital receiving agent. Whether noted or concealed, damage MUST be reported to the carrier immediately upon discovery, or in any event, within 14 days after receipt, and the contents and containers held for inspection by the carrier. A transportation company will not pay a claim for damage if an inspection is not requested within this 14 day period.

CERTIFIED ELECTRICAL CONTRACTOR STATEMENT - FOR USA ONLY

All electrical Installations that are preliminary to positioning of the equipment at the site prepared for the equipment shall be performed by licensed electrical contractors. Other connections between pieces of electrical equipment, calibrations and testing shall be performed by qualified GE personnel. In performing all electrical work on these products, GE will use its own specially trained field engineers. All of GE's electrical work on these products will comply with the requirements of the applicable electrical codes.

The purchaser of GE equipment shall only utilize qualified personnel (i.e., GE's field engineers, personnel of third-party service companies with equivalent training, or licensed electricians) to perform electrical servicing on the equipment.

OMISSIONS & ERRORS

If there are any omissions, errors or suggestions for improving this documentation, please contact the GE Global Documentation Group with specific information listing the system type, manual title, part number, revision number, page number and suggestion details.

Mail the information to:

Service Documentation, GE Medical Systems (China) Co., Ltd. No.19 Changjiang Road WuXi National Hi-Tech Development Zone Jiangsu, P.R China 214028 TEL: +86 510 85225888; FAX: +86 510 85226688

GE employees should use TrackWise to report service documentation issues. These issues will then be in the internal problem reporting tool and communicated to the writer.

SERVICE SAFETY CONSIDERATIONS

DANGER DANGEROUS VOLTAGES, CAPABLE OF CAUSING DEATH, ARE PRESENT IN THIS EQUIPMENT. USE EXTREME CAUTION WHEN HANDLING, TESTING AND ADJUSTING.

WARNING Use all Personal Protection Equipment (PPE) such as gloves, safety shoes, safety glasses, and kneeling pad, to reduce the risk of injury.

For a complete review of all safety requirements, see the Chapter 1, Safety Considerations section in the Service Manual.

LEGAL NOTES

The contents of this publication may not be copied or duplicated in any form, in whole or in part, without prior written permission of GE.

GE may revise this publication from time to time without written notice.

PROPRIETARY TO GE

Permission to use this Advanced Service Software and related documentation (herein called the material) by persons other than GE employees is provided only under an Advanced Service Package License relating specifically to this Proprietary Material. This is a different agreement from the one under which operating and basic service software is licensed. A license to use operating or basic service software does not extend to or cover this software or related documentation.

If you are a GE employee or a customer who has entered into such a license agreement with GE to use this proprietary software, you are authorized to use this Material according to the conditions stated in your license agreement.

However, you do not have the permission of GE to alter, decompose or reverse-assemble the software, and unless you are a GE employee, you MAY NOT COPY the Material. The Material is protected by Copyright and Trade Secret laws; the violation of which can result in civil damages and criminal prosecution.

If you are not party to such a license agreement or a GE Employee, you must exit this Material now.

TRADEMARKS

All products and their name brands are trademarks of their respective holders.

COPYRIGHTS

All Material Copyright© 2014 by General Electric Company Inc. All Rights Reserved.

Revision	Date	Reason for change
1	2014/03/20	Initial Release
2	2014/07/10	Update the CRU list
3	2014/12/23	Update the CRU list

List of Effected Pages (LOEP)

Pages	Revision	Pages	Revision	Pages	Revision
Title Page	3	Chapter 3 - Installation pages 3-1 to 3-28	3	Chapter 8 - Replacement Procedures pages 8-1 to 8-6	3
Important Precautions pages i to x	3	Chapter 4 - Functional Checks pages 4-1 to 4-32	3	Chapter 9 - Replacement Parts pages 9-1 to 9-10	3
Table of Contents pages i to xii	3	Chapter 5 - Theory pages 5-1 to 5-4	3	Chapter 10 - Periodic Maintenance pages 10-1 to 10-18	3
Chapter 1 - Introduction pages 1-1 to 1-17	3	Chapter 6 - Service Adjustments pages 6-1 to 6-2	3	Index pages I to II	3
Chapter 2 - Pre- Installation pages 2-1 to 2-12	3	Chapter 7 - Diagnostics/ Troubleshooting pages 7-1 to 7-10	3	Back Cover	3

Table of Contents

CHAPTER 1 Introduction

Overview	
Purpose of Chapter 1	
Purpose of Service Manual1 - 1	
Typical Users of the Basic Service Manual	
Purpose of Operator Manual(s)1 - 2	
LOGIQ V5/LOGIQ V3 Models Covered by this Manual	
Peripheral List	
Important Conventions 1 5	
Conventions Lead in Book	
Conventions Used in Book	
Safety Considerations	0
Introduction	0
Human Safety	0
Mechanical Safety 1 - 10	0
Electrical Safety	1
Label Locations	1
Dangerous Procedure Warnings1 - 12	2
Returning/Shipping Probes and Repair Parts	3
Lockout/Tagout (LOTO) requirements	л
	-
EMC, EMI, and ESD	5
Electromagnetic Compatibility (EMC)1 - 1	5
CE Compliance	5
Electrostatic Discharge (ESD) Prevention	5
Customer Assistance	G
Contact Information	0
System Manufacturer 1 1	7
$\frac{1}{1} = 1^{-1}$	' 7
	1

CHAPTER 2 Site Preparations

Overview	
General Console Requirements. 2 - 2 Console Environmental Requirements 2 - 2 Electrical Requirements 2 - 2 EMI Limitations 2 - 4	<u>}</u>
Scan Probe Environmental Requirements	;
Purchaser Responsibilities	;
Minimal floor plan suggestion	;
Recommended floor plan suggestion) 0 1

CHAPTER 3 System Setup

Overview.3 - 1Purpose of Chapter 33 - 1Setup Reminders3 - 2Average Installation Time3 - 2Installation Warnings3 - 3Safety Reminders3 - 4	
Receiving and Unpacking the Equipment. 3 - 5 Moving into Position 3 - 9 Product Locator Installation Card 3 - 9	
Preparing for Installation. 3 - 1 Verify Customer Order 3 - 1 Physical Inspection 3 - 1 EMI Protection 3 - 1	0 0 0 0
Completing the Installation3 - 1Power On / Boot Up3 - 1Power Off/ Shutdown3 - 1Transducer Connection3 - 1	1 1 2 2
System Configuration3 - 1System Specifications3 - 1Electrical Specifications3 - 1On-Board Optional Peripherals3 - 1Connecting Cables3 - 1Peripherals/Accessories Connector Panel3 - 1Available Probes3 - 2	3 3 3 4 4
Software/Option Configuration	4
Connectivity Installation Worksheet 3 - 2	5
Loading Base Image Software 3 - 2 Software Version check out 3 - 2 Functional Check-out 3 - 2	6 7 7
Paperwork 3 - 20 Product Locator Installation 3 - 20 User Manual(s) 3 - 20	8 8 8

CHAPTER 4 Functional Checks

Overview
Purpose for Chapter 4
Required Equipment
General Procedure
Power On/Boot Up
Power Off/ Shutdown
Archiving and Loading Presets
Adjusting the Display Monitor
System Features
B Mode Checks
M Mode Controls
Doppler Mode Checks
Color Flow Mode Checks
Basic Measurements
Probe/Connectors Usage 4 - 23
Using Cine
Backup and Restore Database, Preset Configurations and Images 4 - 25
Software Configuration Checks 4 - 32
Peripheral Checks

CHAPTER 5 Components and Functions (Theory)

Overview Purpos	se of Chapter	 5.	 	 	 	 	 	•	 	 	 		 		 • •	 • • • •	•••	 •	5 - 1 5 - 1
Block Diagrar Systen	n n Diagram	 	 	 	 	 	 	•	 	 	 	•	 		 •	 	•	 •	5 - 2 5 - 2
Common Ser Introdu	vice Platform		· · · ·	•••	•••	 •••	 	•	 	· ·	•••	•	 	•	 	 •••		 •	5 - 3 5 - 3

CHAPTER 6 Service Adjustments

Overview	6 ·	- 1
Purpose of this chapter 6	6 -	- 1
Ionitor Adjustments	6 ·	- 2
Adjustments Procedures	6 ·	- 2

CHAPTER 7 Diagnostics/Troubleshooting

Overview	7 - 1 7 - 1
Gathering Trouble Data Overview Collect Vital System Information Collect a Trouble Image with Logs	7 - 2 7 - 2 7 - 2 7 - 3
Screen Captures. Check and Record the Store Key Function Setting the Store Key to Screen Capture Capturing a Screen Reset the Store Key to Customer's Functionality	7 - 4 7 - 4 7 - 4 7 - 5 7 - 6
Common Diagnostics	7 - 7 7 - 7
Network Configuration	7 - 8 7 - 8

CHAPTER 8 Replacement Procedures

Overview
Purpose of Chapter 8
DISASSEMBLY/RE-ASSEMBLY8 - 2
Warning and Caution
Returning/Shipping for repairs8 - 2
Footrest
Loading Base Image Software

CHAPTER 9 Renewal Parts

Overview	9 - 1
Purpose of Chapter 9	9 - 1
List of Abbreviations	9 - 1
Renewal Parts Lists	9 - 2
Equipment Models Covered in this Chapter	9 - 2
Operator Console Assy	9 - 3
CRU Parts List	9 - 4
Manuals	9 - 5
Peripheral	9 - 7
Probes	9 - 9

CHAPTER 10 Care & Maintenance

Overview
Periodic Maintenance Inspections Periodic Maintenance Inspections
Purpose of Chapter 10
Why do Maintenance
Keeping Records
Quality Assurance
Maintenance Task Schedule
How often should care & maintenance tasks be performed?
Tools Required
Standard GE Tool Kit
Special Tools, Supplies and Equipment
Input Power
Cleaning
Physical Inspection
Outlet Test -Wiring Arrangement - USA & Canada
Grounding Continuity
Chassis Leakage Current Test
Isolated Patient Lead (Source) Leakage–Lead to Lead
Isolated Patient Lead (Sink) Leakage-Isolation Test
Probe Leakage Current Test 10 - 13
When There's Too Much Leakage Current

Chapter 1 Introduction

Section 1-1 Overview

1-1-1 Purpose of Chapter 1

This chapter describes important issues related to safely servicing the LOGIQ V5/LOGIQ V3. The service provider must read and understand all the information presented in this manual before installing or servicing a unit.

Table 1-1	Contents i	in Chapter	1
	OOTICOTICO I		

Section	Description	Page Number
1-1	Overview	1-1
1-2	Important Conventions	1-5
1-3	Safety Considerations	1-10
1-4	Lockout/Tagout (LOTO) requirements	1-14
1-5	EMC, EMI, and ESD	1-15
1-6	Customer Assistance	1-16

1-1-2 Purpose of Service Manual

This Service Manual provides installation and service information for the LOGIQ V5/LOGIQ V3 and contains the following chapters:

- 1.) **Chapter 1 Introduction:** Contains a content summary and warnings.
- 2.) **Chapter 2 Site Preparations:** Contains pre-installation requirements for the LOGIQ V5/LOGIQ V3.
- 3.) Chapter 3 System Setup: Contains installation procedures.
- 4.) **Chapter 4 Functional Checks:** Contains functional checks that are recommended as part of the installation, or as required during servicing and periodic maintenance.
- 5.) Chapter 5 Components and Functions (Theory): Contains block diagrams and functional explanations of the electronics.
- 6.) Chapter 6 Service Adjustments: Contains instructions on how to make available adjustments to the LOGIQ V5/LOGIQ V3.
- 7.) Chapter 7 Diagnostics/Troubleshooting: Provides procedures for running diagnostic or related routines for the LOGIQ V5/LOGIQ V3.
- 8.) **Chapter 8 Replacement Procedures:** Provides disassembly procedures and reassembly procedures for all changeable Customer Replaceable Units (CRU).

- 9.) Chapter 9 Renewal Parts: Contains a complete list of field replaceable parts for the LOGIQ V5/ LOGIQ V3.
- 10.) Chapter 10 Care & Maintenance: Provides periodic maintenance procedures for the LOGIQ V5/ LOGIQ V3.

1-1-3 Typical Users of the Basic Service Manual

- Service Personnel (installation, maintenance, etc.).
- Hospital's Service Personnel
- Contractors (Some parts of Chapter 2 Pre-Installation)

1-1-4 Purpose of Operator Manual(s)

The Operator Manual(s) should be fully read and understood before operating the LOGIQ V5/LOGIQ V3 and also kept near the unit for quick reference.

NOTE: Probe information displayed on screen does not necessarily reflect the probes available on your ultrasound system. Please refer to the probe list for available probes and features.

1-1-5 LOGIQ V5/LOGIQ V3 Models Covered by this Manual

Table 1-2LOGIQ V Model Designations

Part Number	Description
5488781	LOGIQ V Color Console (2 Probe Ports)
5478038	LOGIQ V BW Console

1-1-6 Peripheral List

т

Table 1-3 LOGIQ V Model Designations

ltem	Part Name	Part Number	Qty	FRU
	Pi	rinters		•
1000	Sony UP-D897 Chinese kit	5151262	1	N
1000A	Sony UP-D897 USA kit	5151259	1	N
1000B	Sony UP-D897 European kit	5151261	1	N
1000C	Sony UP-D897 Japanese kit	5151263	1	N
1001	Sony UP-D25MD USA kit	5398062	1	N
1001A	Sony UP-D25MD European kit	5398063	1	N
1001B	Sony UP-D25MD Japanese kit	5398064	1	N
1001C	Sony UP-D25MD Chinese kit	5398061	1	N
1002	Sony UP-D711MD with Paper kit	5494719	1	N
1002A	Sony UP-D711MD Paper 1 roll	5494718	1	N
1002B	Sony UP-D711MD Printer Paper 10 rolls	5543548	1	N
1003	HP Officejet 100 Chinese kit	5426594	1	N
1003A	HP Officejet 100 European kit	5426595	1	N
1003A	HP Officejet 100 Japanese kit	5426596	1	N
1003B	HP Officejet 100 USA kit	5426597	1	N
1004	Sony UP-D898MD USA kit	5151259-2	1	N
1004A	Sony UP-D898MD European kit	5151261-2	1	N
1004B	Sony UP-D898MD China kit	5151262-2	1	N
1004C	Sony UP-D898MD Japan kit	5151263-2	1	N
1004D	Sony UP-D898MD Brazil kit	5495509-2	1	N
1005	HP Officejet Pro 8100	NA	1	N
1006	Sony UP-D898MD Printer shelf option kit	5599283	1	N
1007	Sony UP-D711MD Printer shelf option kit	5484791	1	N
Video Converter				
1008	Video Converter Module	5534825	1	N
DVD-RW				
1009	LITEON eUAU108	5454614-2	1	N
Footswitch				
1010	MKF 2-MED GP26 (IPx8)	5151236	1	N
1011	FSU-1000 (IPx8)	5338419	1	N
USB Stick				
1012	SanDisk CRUZER 4G	5395754	1	N

Table 1-3 LOGIQ V Model Designations

ltem	Part Name	Part Number	Qty	FRU
1013	1TB mobile USB HDD	5434317-3	1	N
	Bic	psy Kit		
1014	E8C-RS biopsy kit	E8385NA	1	N
1015	E8C-RS reusable biopsy kit	2398164	1	N
1016	4C-RS biopsy kit	5160703	1	N
1017	L6-12-RS biopsy kit	5176499	1	N
1018	3Sc-RS biopsy kit	5329137	1	N
System and Application Software USB				
1019	LOGIQ V Series R1.0.0 System and Application Software USB	5501727	1	Y
1020	LOGIQ V Series R1.0.1 System and Application Software USB	5501727-2S	1	Y
1021	LOGIQ V Series R1.0.2 System and Application Software USB	5501727-3S	1	Y
1022	LOGIQ V Series R1.0.3 System and Application Software USB	5501727-4S	1	Y

Section 1-2 Important Conventions

1-2-1 Conventions Used in Book

lcons

Pictures, or icons, are used wherever they reinforce the printed message. The icons, labels and conventions used on the product and in the service information are described in this chapter.

Safety Precaution Messages

Various levels of safety precaution messages may be found on the equipment and in the service information. The different levels of concern are identified by a flag word that precedes the precautionary message. Known or potential hazards are labeled in one of following ways:

DANGER DANGER IS USED TO INDICATE THE PRESENCE OF A HAZARD THAT WILL CAUSE SEVERE PERSONAL INJURY OR DEATH IF THE INSTRUCTIONS ARE IGNORED.

- WARNING WARNING IS USED TO INDICATE THE PRESENCE OF A HAZARD THAT CAN CAUSE SEVERE PERSONAL INJURY AND PROPERTY DAMAGE IF INSTRUCTIONS ARE IGNORED.
- **CAUTION** Caution is used to indicate the presence of a hazard that will or can cause minor personal injury and property damage if instructions are ignored.
- **NOTICE Equipment Damage Possible**

Notice is used when a hazard is present that can cause property damage but has absolutely no personal injury risk.

Example: Disk drive will crash.

NOTE: Notes provide important information about an item or a procedure. Information contained in a NOTE can often save you time or effort.

1-2-2 Standard Hazard Icons

Important information will always be preceded by the exclamation point contained within a triangle, as seen throughout this chapter. In addition to text, several different graphical icons (symbols) may be used to make you aware of specific types of hazards that could cause harm.

Table 1-4 Standard Hazard Icons

ELECTRICAL	MECHANICAL	RADIATION
4	3	
LASER	HEAT	PINCH
LASER LIGHT		

Other hazard icons make you aware of specific procedures that should be followed.

	Table 1-5	Standard Icons	Indicating a Special	Procedure Be Used
--	-----------	----------------	-----------------------------	--------------------------

AVOID STATIC ELECTRICITY	TAG AND LOCK OUT	WEAR EYE PROTECTION
		EYE PROTECTION

1-2-3 Product Icons

The following table describes the purpose and location of safety labels and other important information provided on the equipment.

LABEL/SYMBOL	PURPOSE/MEANING	LOCATION
Label/Icon	Purpose/Meaning	Location
Identification and Rating Plate	 Manufacture's name and address Date of manufacture Model and serial numbers Electrical ratings (Volts, Amps, phase, and frequency) 	Refer to User manual for detailed information.
Type/Class Label	Used to indicate the degree of safety or protection.	
IP Code (IPX8) IPX8: FSU-1000, MKF 2-MED GP26	Indicates the degree of protection provided by the enclosure per IEC60 529.	Bottom of footswitch
Ŕ	Type BF Applied Part (man in the box) symbol is in accordance with IEC 60878-02-03.	Beside the probe connector
	General Warning	Various
Â	"CAUTION" - Dangerous voltage" (the lightning flash with arrowhead) is used to indicate electric shock hazards.	Various
ዋ	Indicates the power on and power off position of the power switch. CAUTION: This Power Switch DOES NOT ISOLATE Mains Supply.	See the Console Overview section for location information.
(L)	"Protective Earth" indicates the protective earth (grounding) terminal.	Inside Console
X	This symbol indicates that waste electrical and electronic equipment must not be disposed of as unsorted municipal waste and must be collected separately. Please contact an authorized representative of the manufacturer for information concerning the decommissioning of your equipment.	Bottom

Table 1-6Product Icons

LABEL/SYMBOL	PURPOSE/MEANING	LOCATION
	Indicates the presence of hazardous substance(s) above the maximum concentration value. Maximum concentration values for electronic information products, as set by the People's Republic of China Electronic Industry Standard SJ/T11364-2006, include the hazardous substances of lead, mercury, hexavalent chromium, cadmium, polybrominated biphenyl (PBB), and polybrominated diphenyl ether (PBDE). "20" indicates the number of years during which the hazardous substance(s) will not leak or mutate so that the use of this product will not result in any severe environmental pollution, bodily injury, or damage to any assets.	Probe and Rear Panel, China Rating Plate
	Do not place any objects on the monitor.	Back of LCD
	Do not force the LCD monitor with your hands.	Back of LCD
	There is a pinch point on the LCD monitor. Take care to avoid injuring hands or fingers when flipping down the LCD monitor.	Back of LCD
	Use the rear handle for horizontal movement only.	Back of LCD
×	Do not push the system.	Back of LCD
	"Consult accompanying documents" is intended to alert the user to refer to the operator manual or other instructions when complete information cannot be provided on the label.	Various
PG	GOST symbol: Russia Regulatory Country Clearance.	Bottom Note: Only after Russian regulatory registration is complete, this label will be located on the console rating plate.

Table 1-6	Product	Icons

LABEL/SYMBOL	PURPOSE/MEANING	LOCATION
(Res	Do not put your whole body weight on the foot-rest holder.	Footrest
Segurança Têranî	INMETRO Certification: TUV Rheinland Brazil	Rating plate
		Note: Only after Brazilian regulatory registration is complete, this label will be located on the console rating plate.
e e	NRTL Listing and Certification Mark is used to designate conformance to nationally recognized product safety standards. The Mark bears the name and/or logo of the testing laboratory, product category, safety standard to which conformity is assessed and a control number.	Rear Panel
R ONLY	United States only Presciption Requirement label	Rating Plate

Section 1-3 Safety Considerations

1-3-1 Introduction

The following safety precautions must be observed during all phases of operation, service and repair of this equipment. Failure to comply with these precautions or with specific warnings elsewhere in this manual, violates safety standards of design, manufacture and intended use of the equipment.

1-3-2 Human Safety

Servicing should be performed by authorized personnel only. Only personnel who have participated in a LOGIQ V5/LOGIQ V3 Training are authorized to service the equipment.

1-3-3 Mechanical Safety

WARNING WHEN THE UNIT IS RAISED FOR A REPAIR OR MOVED ALONG ANY INCLINE, USE EXTREME CAUTION SINCE IT MAY BECOME UNSTABLE AND TIP OVER.

WARNING ULTRASOUND PROBES ARE HIGHLY SENSITIVE MEDICAL INSTRUMENTS THAT CAN EASILY BE DAMAGED BY IMPROPER HANDLING. USE CARE WHEN HANDLING AND PROTECT FROM DAMAGE WHEN NOT IN USE. DO NOT USE A DAMAGED OR DEFECTIVE PROBE. FAILURE TO FOLLOW THESE PRECAUTIONS CAN RESULT IN SERIOUS INJURY AND EQUIPMENT DAMAGE.

WARNING NEVER USE A PROBE THAT HAS FALLEN TO THE FLOOR. EVEN IF IT LOOKS OK, IT MAY BE DAMAGED.

CAUTION The LOGIQ V5/LOGIQ V3 weighs 45 kg or more, depending on installed peripherals, when ready for use. Care must be used when moving it or replacing its parts. Failure to follow the precautions listed could result in injury, uncontrolled motion and costly damage.

ALWAYS:

Be sure the pathway is clear.

Use slow, careful motions.

Use two people when moving on inclines or lifting more than 65 kg (143.3 lbs).

WARNING AFTER UNPLUG POWER CORD, WAIT FOR AT LEAST 20 SECONDS FOR CAPACITORS TO DISCHARGE AS THERE ARE NO TEST POINTS TO VERIFY ISOLATION.

1-3-3 Mechanical Safety (cont'd)

NOTE: Special care should be taken when transporting the unit in a vehicle:

- Secure the unit in an upright position.
- Lock the wheels (brake)

1-3-4 Electrical Safety

To minimize shock hazard, the equipment chassis must be connected to an electrical ground. The system is equipped with a three-conductor AC power cable. This must be plugged into an approved electrical outlet with safety ground. The power outlet used for this equipment should not be shared with other types of equipment.

Both the system power cable and the power connector meet international electrical standards.

WARNING DO NOT SERVICE OR DISASSEMBLE PARTS UNDER FRU UNIT LEVEL AT ANY CIRCUMSTANCES.

1-3-5 Label Locations

Refer to User Guide for label location information.

1-3-6 Dangerous Procedure Warnings

Warnings, such as the examples below, precede potentially dangerous procedures throughout this manual. Instructions contained in the warnings must be followed.

DANGER
 DANGEROUS VOLTAGES, CAPABLE OF CAUSING DEATH, ARE PRESENT
 IN THIS EQUIPMENT. USE EXTREME CAUTION WHEN HANDLING, TESTING
 AND ADJUSTING.

	WARNING	EXPLOSION WARNING DO NOT OPERATE THE EQUIPMENT IN AN EXPLOSIVE ATMOSPHERE. OPERATION OF ANY ELECTRICAL EQUIPMENT IN SUCH AN ENVIRONMENT CONSTITUTES A DEFINITE SAFETY HAZARD.
Â	WARNING	DO NOT SUBSTITUTE PARTS OR MODIFY EQUIPMENT BECAUSE OF THE DANGER OF INTRODUCING ADDITIONAL HAZARDS, DO NOT INSTALL SUBSTITUTE PARTS OR PERFORM ANY UNAUTHORIZED MODIFICATION OF THE EQUIPMENT.
Â	WARNING	SHUT DOWN FORCEDLY OR PLUG IN/OUT ACDC INVALID MAY CAUSE THE DAMAGE OF SYSTEM FILES.
Â	WARNING	AFTER UNPLUG POWER CORD, WAIT FOR AT LEAST 20 SECONDS FOR CAPACITORS TO DISCHARGE AS THERE ARE NO TEST POINTS TO VERIFY ISOLATION.
1-3-7 Returning/Shipping Probes and Repair Parts

Equipment being returned must be clean and free of blood and other infectious substances.

GEMS policy states that body fluids must be properly removed from any part or equipment prior to shipment. GEMS employees, as well as customers, are responsible for ensuring that parts/equipment have been properly decontaminated prior to shipment. Under no circumstance should a part or equipment with visible body fluids be taken or shipped from a clinic or site (for example, body coils or an ultrasound probe).

The purpose of the regulation is to protect employees in the transportation industry, as well as the people who will receive or open this package.

NOTE: The USER/SERVICE staff should dispose all the waste properly as per federal, state, and local waste disposal regulation.

Equipment being returned must be clean and free of blood and other infectious substances.

GE policy states that body fluids must be properly removed from any part or equipment prior to shipment. GE employees, as well as customers, are responsible for ensuring that parts/equipment have been properly decontaminated prior to shipment. Under no circumstance should a part or equipment with visible body fluids be taken or shipped from a clinic or site (for example, body coils or an ultrasound probe).

The purpose of the regulation is to protect employees in the transportation industry, as well as the people who will receive or open this package.

- NOTE: The US Department of Transportation (DOT) has ruled that "items that were saturated and/or dripping with human blood that are now caked with dried blood; or which were used or intended for use in patient care" are "regulated medical waste" for transportation purposes and must be transported as a hazardous material.
- NOTE: The USER/SERVICE staff should dispose all the waste properly as per federal, state, and local waste disposal regulation.

Section 1-4 Lockout/Tagout (LOTO) requirements

Follow OSHA Lockout/Tagout requirements (USA) or local Lockout/Tagout requirements by ensuring you are in total control of the AC power plug at all times during the service process.

To apply Lockout/Tagout:

- 1.) Plan and prepare for shutdown.
- 2.) Shutdown the equipment.
- 3.) Isolate the equipment.
- 4.) Apply Lockout/Tagout Devices.
- 5.) Control all stored and residual energy.
- 6.) Verify isolation.

All potentially hazardous stored or residual energy is relieved.



Section 1-5 EMC, EMI, and ESD

1-5-1 Electromagnetic Compatibility (EMC)

Electromagnetic compatibility describes a level of performance of a device within its electromagnetic environment. This environment consists of the device itself and its surroundings including other equipment, power sources and persons with which the device must interface. Inadequate compatibility results when a susceptible device fails to perform as intended due interference from its environment or when the device produces unacceptable levels of emission to its environment. This interference is often referred to as radio–frequency or electromagnetic interference (RFI/EMI) and can be radiated through space or conducted over interconnecting power of signal cables. In addition to electromagnetic energy, EMC also includes possible effects from electrical fields, magnetic fields, electrostatic discharge and disturbances in the electrical power supply.

1-5-2 CE Compliance

The LOGIQ V5/LOGIQ V3 unit conforms to all applicable conducted and radiated emission limits and to immunity from electrostatic discharge, radiated and conducted RF fields, magnetic fields and power line transient requirements.

For applicable standards refer to the Safety Chapter in the User Guide.

NOTE: For CE Compliance, it is critical that all covers, screws, shielding, gaskets, mesh, clamps, are in good condition, installed tightly without skew or stress. Proper installation following all comments noted in this service manual is required in order to achieve full EMC performance.

1-5-3 Electrostatic Discharge (ESD) Prevention

/ WARNING

DO NOT TOUCH ANY BOARDS WITH INTEGRATED CIRCUITS PRIOR TO TAKING THE NECESSARY ESD PRECAUTIONS:



1.FOLLOW GENERAL GUIDELINES FOR HANDLING OF ELECTROSTATIC SENSITIVE EQUIPMENT.

Section 1-6 Customer Assistance

1-6-1 Contact Information

If this equipment does not work as indicated in this service manual or in the User Manual, or if you require additional assistance, please contact the local distributor or appropriate support resource, as listed below.

Prepare the following information before you call:

- System ID serial number.
- Software version.

LOCATION		PHON	IE NUMBER
USA GE Medical Systems Ultrasound Service Engineering 9900 Innovation Drive Wauwatosa, WI 53226	Service: Service F Application	On-site Parts on Support	1-800-437-1171 1-800-558-2040 1-800-682-5327 or 1-262-524-5698
Canada			1-800-668-0732
Latin America	Service Applicati	on Support	1-800-321-7937 1-262-524-5698
Europe (OLC- EMEA) GE Ultraschall Deutschland GmbH Beethovenstraße 239 Postfach 11 05 60, D-42655 Solingen Germany	OLC - EN Phone: Fax:	MEA +49 (0)212 2802 - 6 +33 1 3083 1300 +49 (0) 212 2802 - 4	52 431
Online Services Ultrasound Asia Australia China India Japan Korea Singapore	Phone:	+(61) 1-800-647-853 +(86) 800-810-8188 +(91) 1-800-11-4563 +(81) 42-648-2924 +(82) 2620 13585 +(95) 6277-3444	5

Table 1-7 Phone Numbers for Customer Assistance

1-6-2 System Manufacturer

Table 1-8 System Manufacturer

Manufacturer	FAX Number
GE Medical Systems (China) Co., Ltd. No.19, Changjiang Road, Wuxi National Hi-Tech Development Zone, Jiangsu, P.R. China 214028	TEL: +86 510-85225888 FAX: +86 510-85226688

1-6-3 Factory Site

Table 1-9 Factory Site

Factory Site	FAX Number
GE Medical Systems (China) Co., Ltd. No.19, Changjiang Road, Wuxi National Hi-Tech Development Zone, Jiangsu, P.R. China 214028	TEL: +86 510-85225888 FAX: +86 510-85226688

This page was intentionally left blank.

Chapter 2 Site Preparations

Section 2-1 Overview

2-1-1 Purpose of chapter 2

This chapter provides the information required to plan and prepare for the installation of LOGIQ V5/ LOGIQ V3. Included are descriptions of the facility and electrical needs to be met by the purchaser of the unit.

Table 2-1	Contents in Chapter 2	
	Contents in Chapter Z	

Section	Description	Page Number
2-1	Overview	2-1
2-2	General Console Requirements	2-2
2-3	Facility Needs	2-6

-

Section 2-2 General Console Requirements

2-2-1 Console Environmental Requirements

Table 2-2 Environmental Requirements for LOGIQ V Series Scanners

	Operational	Storage	Transport
Temperature	10 - 40°C	-5 - 50 °C	-5 - 50 ºC
	50- 104 °F	23 - 122°F	23 - 122ºF
Humidity	30 - 80%	10 - 90%	10 - 90%
	non-condensing	non-condensing	non-condensing
Pressure	700 - 1060hPa	700 - 1060hPa	700 - 1060hPa

NOTE: Temperature in degrees C. Conversion to Degrees F = (Degrees C * 9/5) + 32.

2-2-1-1 Lighting

Bright light is needed for system installation, updates and repairs. However, operator and patient comfort may be optimized if the room light is subdued and indirect. Therefore a combination lighting system (dim/bright) is recommended. Keep in mind that lighting controls and diameters can be a source of EMI which could degrade image quality. These controls should be selected to minimize possible interface.

2-2-2 Electrical Requirements

NOTE: GE Healthcare requires a dedicated power and ground for the proper operation of its Ultrasound equipment. This dedicated power shall originate at the last distribution panel before the system.

Sites with a mains power system with defined Neutral and Live:

The dedicated line shall consist of one phase, a neutral (not shared with any other circuit), and a full size ground wire from the distribution panel to the Ultrasound outlet.

Sites with a mains power system without a defined Neutral:

The dedicated line shall consist of one phase (two lines), not shared with any other circuit, and a full size ground wire from the distribution panel to the Ultrasound outlet.

NOTE: Please note that image artifacts can occur, if at any time within the facility, the ground from the main facility's incoming power source to the Ultrasound unit is only a conduit.

2-2-2Electrical Requirements (cont'd)

2-2-2-1 LOGIQ V5/LOGIQ V3 Power Requirements

The following power line parameters should be monitored for one week before installation. We recommend that you use an analyzer Dranetz Model 606-3 or Dranetz Model 626:

Table 2-3 Electrical Specifications for LOGIQ V5/LOGIQ V3

PARAMETER	AREA	LIMITS
Voltage Range	100-240V~	300VA
Power	All applications	MAX. 300 VA
Line Frequency	All applications	50/60Hz
Power Transients	All applications	Less than 25% of nominal peak voltage for less than 1 millisecond for any type of transient, including line frequency, synchronous, asynchronous, or aperiodic transients.
Decaying Oscillation	All applications	Less than 15% of peak voltage for less than 1 millisecond.

2-2-2-2 Inrush Current

Inrush Current is not a factor to consider due to the inrush current limiting properties of the power supplies.

2-2-2-3 Site Circuit Breaker

It is recommended that the branch circuit breaker for the machine be ready accessible.

CAUTION POWER OUTAGE MAY OCCURE.

The LOGIQ V5/LOGIQ V3 requires a dedicated single branch circuit. To avoid circuit overload and possible loss of critical care equipment, make sure you DO NOT have any other equipment operating on the same circuit.

2-2-2-4 Site Power Outlets

A desiccated AC power outlet must be within reach of the unit without extension cords. Other outlets adequate for the external peripherals, medical and test equipment needed to support this unit must also be present within 1 m (3.2 ft.) of the unit. Electrical installation must meet all current local, state, and national electrical codes.

2-2-2-5 Unit Power Plug

If the unit arrives without the power plug, or with the wrong plug, you must contact your GE dealer or the installation engineer must supply what is locally required.

2-2-2-6 Power Stability Requirements

Voltage drop-out

Max 10 ms.

Power Transients

(All applications)

Less than 25% of nominal peak voltage for less than 1 millisecond for any type of transient, including line frequency, synchronous, asynchronous, or aperiodic transients.

2-2-3 EMI Limitations

Ultrasound machines are susceptible to Electromagnetic Interference (EMI) from radio frequencies, magnetic fields, and transient in the air wiring. They also generate EMI. The LOGIQ V5/LOGIQ V3 complies with limits as stated on the EMC label. However there is no guarantee that interface will not occur in a particular installation.

Possible EMI sources should be identified before the unit is installed.

Electrical and electronic equipment may produce EMI unintentionally as the result of defect.

These sources include:

- medical lasers,
- scanners,
- cauterizing guns,
- computers,
- monitors,
- fans,
- gel warmers,
- microwave ovens,
- light dimmers,
- portable phones.

The presence of broadcast station or broadcast van may also cause interference. See for EMI Prevention tips.

See Table 2-4 for EMI Prevention tips.

Table 2-4 LIVIL Prevention/abatement	Table 2-4	EMI Prevention/abatement
--------------------------------------	-----------	--------------------------

EMI Rule	Details
Be aware of RF sources	Keep the unit at least 5 meters or 15 feet away from other EMI sources. Special shielding may be required to eliminate interference problems caused by high frequency, high powered radio or video broadcast signals.
Ground the unit	Poor grounding is the most likely reason a unit will have noisy images. Check grounding of the power cord and power outlet.
Replace all screws, RF gaskets, covers, cores	After you finish repairing or updating the system, replace all covers and tighten all screws. Any cable with an external connection requires a magnet wrap at each end. Install the shield over the front of card cage. Loose or missing covers or RF gaskets allow radio frequencies to interface with the ultrasound signals.
Replace broken RF gaskets	If more than 20% or a pair of fingers on the RF gaskets are broken, replace the gaskets. Do not turn on the unit until any loose metallic part is removed.
Do not place labels where RF gaskets touch metal	Never place a label where RF gaskets meet the unit. Otherwise, the gap created will permit RF leakage. Or, if a label has been found in such a position, move the label.
Use GE specified harnesses and peripherals	The interconnect cables are grounded and require ferrite beads and other shielding. Also, cable length, material, and routing are all important; do not change from what is specified.
Take care with cellular phones	Cellular phones may transmit a 5 V/m signal; that could cause image artifacts.
Properly dress peripheral cables	Do not allow cables to lie across the top of the card cage or hang out of the peripheral bays. Loop the excess length for peripheral cables inside the peripheral bays. Attach the monitor cables to the frame.

2-2-4 Scan Probe Environmental Requirements

Operation: 10° to 40° C

Storage: -5° to 50° C

NOTE: Temperature in degrees C. Conversion to Degrees F = (Degrees C * (9/5) + 32).

NOTICE SYSTEMS AND ELECTRONIC PROBES ARE DESIGNED FOR STORAGE TEMPERATURES OF -10 TO + 60 degrees C. WHEN EXPOSED TO LARGE TEMPERATURE VARIATIONS, THE PRODUCT SHOULD BE KEPT IN ROOM TEMPERATURE FOR 10 HOURS BEFORE USE.

Section 2-3 Facility Needs

2-3-1 Purchaser Responsibilities

The work and materials needed to prepare the site is the responsibility of the purchaser. Delay, confusion, and waste of manpower can be avoided by completing pre installation work before delivery. User the Pre Installation checklist to verify that all needed steps have been taken, Purchaser reasonability includes:

- Procuring the materials required.
- Completing the preparations before delivery of the ultrasound system.
- Paying the costs for any alternations and modifications not specifically provided in the sales contract.

NOTE: All electrical installation that are preliminary to the positioning of the equipment at the site prepared for the equipment must be performed by licensed electrical contractors. Other connections between pieces of electrical equipment, products involved (and the accompanying electrical installations) are highly sophisticated and special engineering competence is required. All electrical work on these product must comply with the requirements of applicable electrical codes. The purchaser of GE equipment must only utilize qualified personnel to perform electrical servicing on the equipment.

The desire to use a non-listed or customer provided product or to place an approved product further from the system than the interface kit allows presents challenges to the installation team. To avoid delays during installation, such variances should be made known to the individuals or group performing the installation at the earliest possible date (preferable prior to purchase).

The ultrasound suite must be clean proof to delivery of the machine. Carpet is not recommended because it collects dust and creates static. Potential sources of EMI (electromagnetic interference) should also be investigated before delivery. Dirt, static, and EMI can negatively impact system.

2 - 6

2-3-2 Required Features

NOTE: GE Medical Systems requires a dedicated power and ground for the proper operation of its Ultrasound equipment. This dedicated power shall originate at the last distribution panel before the system.

Sites with a mains power system with defined Neutral and Live:

The dedicated line shall consist of one phase, a neutral (not shared with any other circuit), and a full size ground wire from the distribution panel to the Ultrasound outlet.

Sites with a mains power system without a defined Neutral:

The dedicated line shall consist of one phase (two lines), not shared with any other circuit, and a full size ground wire from the distribution panel to the Ultrasound outlet.

Please note that image artifacts can occur, if at any time within the facility, the ground from the main facility's incoming power source to the Ultrasound unit is only a conduit.

- Dedicated single branch power outlet of adequate amperage meeting all local and national codes which is located less than 2.5 m (8 ft.) from the unit's proposed location
- Door opening is at least 76 cm (30 in) wide
- Proposed location for unit is at least 0.3 m (1 ft.) from the wall for cooling
- Power outlet and place for any external peripheral are within 2 m (6.5 ft.) of each other with peripheral within 1 m of the unit to connect cables.

2-3-3 Minimal floor plan suggestion

CSI 8x10



- 1. Door at least 762 mm (30 inches)
- 2. Film Viewer
- 3. Counter Top, Sink with hot and cold water and Supplies Storage
- 4. Linen Supply
- 5. Probes/Supplies

- 6. Examination Table 1930 x 610 mm
- (76 x 24 inches)7. Footswitch
- 8. Stool
- 9. Ultrasound system
- 10. External Peripherals
- 11. Dedicated Power Outlet Circuit Breaker protected and easily accessible
- Figure 2-1. Minimal floor plan, 2.5 m x 3 m (8 by 10 foot)

Scale:

Each square equals one square foot (app. 31 x 31 cm)

- 12. Network Interface
- 13. 457 mm (18 inches) distance of Ultrasound system from wall or objects
- 14. GE Cabinet for Software and Manuals

2-3-4 Recommended floor plan suggestion

Recommended standard floor plan and a minimal floor plan for ultrasound equipment:



Scale: Each square equals one square foot (app. 31 x 31 cm)

- 1. Secretaries or Doctors Desk
- 2. File Cabinet
- 3. Film Viewer
- 4. Counter Top
- 5. Counter Top and Sink with hot and cold water
- 6. Overhead Lights Dimmer Dual Level Lighting (bright and dim)
- 7. Emergency Oxygen

- 8. Suction Line
- 9. Ultrasound system
- 10. Dedicated Power Outlet Circuit Breaker protected and easily accessible
- 11. Network Interface
- 12. 457 mm (18 inches) distance of Ultrasound system from wall or objects
- 13. Stool

- 14. Footswitch
- 15. Storage for Linens and Equipment
- 16. Examination Table 1930 x 610 mm (76 x 24 inches)
- 17. Lavatory and Dressing Room
- 18. Door at least 762 mm (30 inches)

Suggested floor plan, Ultrasound system, and EchoPAC PC in same room 2-3-5



1. EchoPAC PC workstation parts 2. UPS Ethernet network wall outlet

3.

- 4. 3x mains power outlets Hot and Cold water 5.
- 6. Dedicated mains power outlet

Figure 2-2. Suggested Room with EchoPAC PC workstation and Ultrasound Scanner

2-3-6 Networking Pre-installation Requirements

2-3-6-1 Stand Alone Scanner (without Network Connection) None.

2-3-6-2 Scanner Connected to Hospital's Network

Supported networks:

Wire LAN

2-3-6-3 Purpose of DICOM Network Function

DICOM services provide the operator with clinically useful features for moving images and patient information over a hospital network. Examples of DICOM services include the transfer of images to workstations for viewing or transferring images to remote printers. As an added benefit, transferring images in this manner frees up the on-board monitor and peripherals, enabling viewing to be done while scanning continues. With DICOM, images can be archived, stored, and retrieved faster, easier, and at a lower cost.

2-3-6-4 DICOM Option Pre-installation Requirements

To configure the LOGIQ V5/LOGIQ V3 to work with other network connections, the site's network administrator must provide some necessary information.

Information must include:

- A host name, local port number, AE Title, IP address and Net Mask for the LOGIQ V5/LOGIQ V3.
- The IP addresses for the default gateway and other routers at the site for ROUTING INFORMATION.
- The host name, IP address, port and AE Title for each device the site wants connected to the LOGIQ V5/LOGIQ V3 for DICOM APPLICATION INFORMATION. A field for the make (manufacturer) and the revision of the device, is also included. This information may be useful for solving errors.

2-3-6-4 DICOM Option Pre-installation Requirements (cont'd)

LOGIQ V Host Nan AE Title ROUTING	5/ ne INFORMATION ROUTER1 ROUTER2 ROUTER3	Loc Destinatio IP Address	al Port n ses 	IP Address Net Mask Default	GATEWAY IP Addresses
DICOM A	PPLICATION INFORMA	TION				
	NAME	MAKE/REVISION			DRESSES	PORT
Store 1						
Store 2						
Store 3						
Store 4	· · · · · · · · · · · · · · · · · · ·					
Store 5						
Store 6						
Worklist				·····		
Storage Commit						
MPPS				·····		

Figure 2-2 Worksheet for DICOM Network Information

Chapter 3 System Setup

Section 3-1 Overview

3-1-1 Purpose of Chapter 3

This chapter contains information needed to install the unit. Included are references to a procedure that describes how to receive and unpack the equipment and how to file a damage or loss claim. How to prepare the facility and unit of the actual installation, and how to check and test the unit and external peripherals for electrical safety are included in this procedure. Also included LOGIQ V5/LOGIQ V3 in this section are guidelines for transporting the unit to a new site.

Section	Description	Page Number
3-1	Overview	3-1
3-2	Setup Reminders	3-2
3-3	Receiving and Unpacking the Equipment	3-5
3-4	Preparing for Installation	3-10
3-5	Completing the Installation	3-11
3-6	System Configuration	3-13
3-7	Software/Option Configuration	3-7
3-8	Connectivity Installation Worksheet	3-25
3-9	Loading Base Image Software	3-26
3-10	Software Version check out	3-27
3-11	Paperwork	3-28

Table 3-1 Contents in Chapter 3

Section 3-2Setup Reminders

3-2-1 Average Installation Time

 Table 3-2
 Average Installation Time

Description	Average Installation Time	Comments
Unpacking the scanner	0.5 hour	
Scanner wo/options	0.5 hour	Dependant on the configuration that is required
DICOM Option	0.5 hour	Dependant on the amount of configuration

The LOGIQ V5/LOGIQ V3 has been designed to be installed and checked out by an experienced service technician in approximately 1 hour. LOGIQ V5/LOGIQ V3 consoles with optional equipment may take slightly longer.

3-2-2 Installation Warnings

- 1.) Since the LOGIQ V5/LOGIQ V3 weighs approximately 50 kg without options, preferably two people should unpack it. Two people are also preferable for installing any additional bulky items.
- 2.) There are no operator serviceable components. To prevent shock, do not remove any covers or panels. Should problems or malfunctions occur, unplug the power cord. Only qualified service personnel should carry out servicing and troubleshooting.
- NOTE: For information regarding packing labels, refer to LABELS ON PACKAGE.
 - 3.) After being transported, the unit may be very cold or hot. If this is the case, allow the unit to acclimate before you turn it on. It requires one hour for each 2.5°C increment it's temperature is below 10°C or above 30°C.

CAUTION Equipment damage possibility. Turning the system on without acclimation after arriving at site may cause the system to be damaged.

°C	60	55	50	45	40	35	30	25	20	15	10	5	0	-5	-10	-15	-20	-25	-30	-35	-40
°F	140	131	122	113	104	95	86	77	68	59	50	41	32	23	14	5	-4	-13	-22	-31	-40
hrs	8	6	4	2	0	0	0	0	0	0	0	2	4	6	8	10	12	14	16	18	20

Table 3-3Acclimation Time

3-2-3 Safety Reminders

Â	DANGER	WHEN USING ANY TEST INSTRUMENT THAT IS CAPABLE OF OPENING THE
		AC GROUND LINE (I.E., METER'S GROUND SWITCH IS OPEN), DON'T TOUCH
		THE UNIT!

- **CAUTION** Two people should unpack the unit because of its weight. Two people are required whenever a part weighing 19kg (35 lb.) or more must be lifted.
- **CAUTION** If the unit is very cold or hot, do not turn on its power until it has had a chance to acclimate to its operating environment.
- **CAUTION** To prevent electrical shock, connect the unit to a properly grounded power outlet. Do not use a three to two prong adapter. This defeats safety grounding.
- **CAUTION** Do NOT wear the ESD wrist strap when you work on live circuits and more than 30 V peak is present.

CAUTIONDo not use a 20 Amp to 15 Amp adapter on the 120 Vac unit's power cord. This unit requires
a dedicated 20 A circuit and can have a 15A plug if the on board peripherals do not cause the
unit to draw more than 14.0 amps.

CAUTIONDo not operate this unit unless all board covers and frame panels are securely in place.System performance and cooling require this.

CAUTION OPERATOR MANUAL(S) The User Manual(s) should be fully read and understood before operating the LOGIQ V5/ LOGIQ V3 and kept near the unit for quick reference.



Figure 3-1 Environmental Labels

Section 3-3 Receiving and Unpacking the Equipment

When a new system arrives, check that any components are not damaged and are not in short supply. If shipping damage or shortage occurs, contact the address shown in Chapter 1.

CAUTION Do not lift the unit by the Keyboard. Equipment damage may result.

CAUTION The crate with the LOGIQ V5/LOGIQ V3 weighs approximately 50kg. Be prepared for a sudden shift of weight as the unit is removed from its base (pallet)

Step	Description	Corresponding Graphic
1	Tear the stop open mark.	V Series
2	Cut the two packing straps around the crate. Note: To avoid injury, with one hand holding the strap clasp when cutting the strap.	Les ries
3	Remove the top cover of the shipping box.	

Table 3-4:	Unpacking	the equipment
	• · · · • · · · · · · · · · · · · · · ·	

Step	Description	Corresponding Graphic
4	Open the four plastic locks.	
	Note: Rotate the inside plastic lock counterclockwise to remove it and then remove the outside lock.	
5	Remove the outside shipping box.	
6	Remove the dust bag from the unit.	

Table 3-4: Unpacking the equipment

Step	Description	Corresponding Graphic
7	Remove the clear plastic (wrapped around the LOGIQ V5/LOGIQ V3) from the unit. Note: To avoid damaging the unit, please use a pair of scissors instead of the knife.	
8	Remove the accessory box.	

Table 3-4: Unpacking the equipment

Step	Description	Corresponding Graphic
9	Remove the foams beside the LCD monitor and the control panel.	
10	Cut the packing straps around the four wheels and remove the foams beside the wheels. Note: To aviod injury, lock the wheels before cutting the packing straps.	
11	Unlock the wheels, and then hold the control panel at the front side to move the system until the front two wheels on the ground, and then slowly move the whole system on the ground.	

Table 3-4: Unpacking the equipment

3-3-1 Moving into Position

CAUTION Do not tilt the unit more than 5 degrees to avoid tipping it over.

To avoid injury by tipping over. Set the monitor to the lowest position before moving.

In general, a single adult can move the LOGIQ V5/LOGIQ V3 along an even surface with no steep grades. At least two people should move the machine when large humps, grooves, or grades will be encountered. (It is better to pull from the rear rather than push from the front of the unit). Before moving, store all loose parts in the unit. Wrap transducers in soft cloth or foam to prevent damage.

Although LOGIQ V5/LOGIQ V3 is a mobile machine, two people should move it over rough surfaces or up and down grades.

3-3-2 Product Locator Installation Card

(Mailing Address	GE Medic Product Lo P.O. Box Milwaukee	al Sy ocato 414 e, Wi	stem or File 5320	s 9 01-0414						
	DESCRIPTION		FDA	MODE	L			REV	SERIAL		
	PREPARE FOR ORDERS THAT	DO NOT			OCP	BS	ORD			DATE (MO-DA-YR)	
	HAVE A LOCATOR INSTALLATION	REPORT			DISTCOUNTRY	ROOM				EMPLOYEE NO.	
NSA	SYSTEM ID NUMBER				CUSTOMER NO.						
IN TED IN	INSTALLATI	O N			DESTINATION - N	AME AND ADI	ORESS				
N Bd											
LALLATIO											
LSNI										ZIP CODE	

Figure 3-2 Product Locator Installation Card

NOTE: The Product Locator Installation Card shown may not be same as the provided Product Locator card.

Section 3-4 Preparing for Installation

3-4-1 Verify Customer Order

Compare items received by the customer to that which is listed on the delivery order. Report any items that are missing, back ordered or damaged.

3-4-2 Physical Inspection

3-4-2-1 System Voltage Settings

Verify that Docking Cart is set to the correct voltage. The Voltage settings for the LOGIQ V5/LOGIQ V3 is found on a label to the right of the Power switch and External I/O, on the rear of the system.

WARNING Connecting a LOGIQ V5/LOGIQ V3 to the wrong voltage level will most likely destroy it.

3-4-2-2 Video Formats

Check that the video format is set to the locally used video standard, NTSC or PAL.

3-4-3 EMI Protection

This Unit has been designed to minimize the effects of Electro Magnetic Interference (EMI). Many of the covers, shields, and screws are provided primarily to protect the system from image artifacts caused by this interference. For this reason, it is imperative that all covers and hardware are installed and secured before the unit is put into operation.

Section 3-5 Completing the Installation

3-5-1 Power On / Boot Up

NOTE: After turning off a system, wait at least ten seconds before turning it on again. The system may not be able to boot if power is recycled too quickly.

3-5-1-1 Scanner Power On

When power is applied to the scanner, power is distributed to the Cooling Unit, Control Panel, LCD, Peripherals and the Back-end Processor.

3-5-1-2 Turn on the system

Press the Power On/Off switch at the front of the system once.



Figure 3-3 Power On/Off Switch

When the *Power On/Off* switch on the Control Panel is pressed once, the Back-end Processor starts and the software code is distributed to initiate the scanner.

No status messages are displayed during this process.

3-5-2 Power Off/ Shutdown

NOTE: After turning off a system, wait at least ten seconds before turning it on again. The system may not be able to boot if power is recycled too quickly.

3-5-2-1 Back-end Processor Power Down

To power down the system:

- 1.) Press the Power On/Off switch once.
- 2.) The System-Exit window is displayed.

-	Logen Information
Sys	tem Administrator is logged on as ADM
Logon Time	2014.05.08 - 16:16
So	Itware Remote Upgrade Information
Softwa	are Download Service connection failed

Figure 3-4 System-Exit Window

- 3.) Using the Set key, select Shutdown.
- 4.) The shutdown process takes a few seconds and is complete when the power status LED is turned blue.
- 5.) Disconnect the probes. Clean or disinfect all probes as necessary. Store them in their shipping cases to avoid damage.

3-5-2-2 Scanner Shutdown

Disconnect the Mains Power Cable if necessary. For example: Relocating the scanner.

3-5-3 Transducer Connection

- 1.) Plug the probe connector into the probe port, then lock the probe.
- NOTE: Please ensure that the probe latch is in an unlocked position before you connect the probe to the system.
- NOTE: It is not necessary to turn OFF power to connect or disconnect a probe.

Section 3-6 System Configuration

3-6-1 System Specifications

3-6-1-1 Physical Dimensions

The physical dimensions of the LOGIQ V5/LOGIQ V3 unit are summarized in Table 3-5 on page 3-13. The Size of LOGIQ V5/LOGIQ V3.

Table 3-5 Physical Dimensions of LOGIQ V5/LOGIQ V3

Height	Width Depth		Unit
1320±60	420±30 (Wheel to Wheel)	635±30	mm
52±2.3	16.5±1.1	25±1.1	inches

3-6-1-2 Weight

Table 3-6 Weight of LOGIQ V5/LOGIQ V3 With Monitor and Without Other Peripherals

Model	Weight [kg]	Weight [lb]
LOGIQ V5/LOGIQ V3	Approximately 45	Approximately 99.2

3-6-2 Electrical Specifications

Table 3-7 Electrical Specifications for LOGIQ V5/LOGIQ V3

Voltage	Current	Frequency
100 -240 V AC	300VA	50/60Hz

3-6-3 On-Board Optional Peripherals

Table 3-8List of Optional Peripherals

Device	Manufacturer	Model	Video Signal
B/W Printer	SONY	UP-D897MD	USB
B/W Printer	SONY	UP-D898MD	USB
B/W Printer	SONY	UP-D711MD	USB
HP Printer	HP	HP Officejet 100	USB
HP Printer	HP	HP Officejet Pro 8100	USB
USB Memory	SanDisk	SanDisk 4G	USB
1-Pedal Footswitch	Whanam	FSU1000	USB
3-Pedal Footswitch	Whanam	MKF 2-MED GP26	USB
USB Hard Disk	Seagate	USB HDD 500G	USB
Color Printer	SONY	UP-D25MD	USB

3-6-4 **Connecting Cables**

WARNING Equipment damage possibility. Be sure to use the following recommended connecting cables to connect recording devices and a network with LOGIQ V5/LOGIQ V3 console.

Table 3-9 **List of Connecting Cables**

Name	Part No.	Figure	NOTE
USB Cable			For USB Printer

3-6-5 **Peripherals/Accessories Connector Panel**

LOGIQ V5/LOGIQ V3 peripherals and accessories can be properly connected using the side connector panel.

Rear Panel Connector 3-6-5-1



Figure 3-5 Rear Connector Panel

NOTE: The AC printer can be connected to the isolated Printer USB port on the rear panel only.

3 - 14

Section 3-3 - Receiving and Unpacking the Equipment

3-6-5Peripherals/Accessories Connector Panel (cont'd)

Table 3-10 Peripheral/Accessory Connector Panel

1	HDMI port	HDMI out
2	Isolated Printer USB port	For AC Printer ONLY
3	USB Ports	USB Ports
4	Ethernet	LAN for InSite Connection (RJ45)
5	Circuit breaker	6.5A
6	AC Inlet	100-240V

3-6-5-2 Pin assignment for each connector

Table 3-11 Pin Assignments of External VGA

Pin No.	Signal	Pin No.	Signal
1	RED	9	NC
2	GREEN	10	NC
3	BLUE	11	NC
4	NC	12	NC
5	NC	13	HSY
6	GND	14	VSY
7	GND	15	NC
8	GND		

Pin No.	Signal	Pin No.	Signal
1	+5 VDC	5	+5 VDC
2	DATA	6	DATA
3	DATA	7	DATA
4	GND	8	GND

Table 3-13 Pin Assignments of Audio

Pin No.	Signal	Pin No.	Signal
1	GND	4	NC
2	L+	5	R+
3	Speaker L	6	Speaker R

Table 3-14 Pin Assignment of S-Video

Pin No.	Output Signal	Pin No.	Output Signal
1	GND	3	Y
2	GND	4	С

Table 3-15 Pin Assignment of Composite Video Out

Pin No.	Output Signal	Pin No.	Output Signal
1	Composite Out	2	GND

3-6-5-3 Connect Peripherals

A.) Connect B/W printer to the system.

Sony UP-D711MD Printer can be properly connected using the STD and DC cables.



Figure 3-6 Connect B/W printer to the system

Sony UP-D711MD is connected to the LOGIQ V5/LOGIQ V3 system via the STD and DC cables under the control panel.



Figure 3-7 STD cable location

3-6-5-3Connect Peripherals (cont'd)

B.) Connect Sony UP-D25MD color printer to the system. Sony UP-D25MD Color Printer can be properly connected using the isolated Printer USB Port. And connect the power cable of Sony UP-D25MD to the wall outlet.



Figure 3-8 Connect color printer to the system
3-6-5-3 Connect Peripherals (cont'd)

C.) Connect the B/W printer (Sony UP-897/Sony UP-D898MD) to the system The B/W printer can be properly connected using the isolated Printer USB Port. And connect the power cable of Sony UP-D897/Sony UP-D898MD to the wall outlet.



Figure 3-9 Connect B/W Printer (Sony UP-897) to the system

D.) Connect HP Officejet 100 printer to the system. HP Officejet 100 Printer can be properly connected using the isolated Printer USB Port. And connect the power cable of HP Officejet 100 to the wall outlet.



Figure 3-10 HP Officejet 100 printer to the system

3-6-5-3Connect Peripherals (cont'd)

- E.) Connect Foot Switch to the system.
 - Foot Switch can be properly connected using USB Ports.





1-Pedal Footswitch

3-Pedal Footswitch

Figure 3-11 Connect Foot Switch to the system

3-6-5-3Connect Peripherals (cont'd)

F.) Connect the USB Memory to the system. The USB Memory can be properly connected using the USB ports.



Figure 3-12 USB Memory Connection

G.) Connect the USB HDD to the system. The USB Harddisk can be properly connected using USB ports.



Figure 3-13 USB Hard Disk Connection

3-6-5-4 Digital Printer Setup

There are two steps to do when setting up a digital printer: 1. follow the procedure below for each printer, then 2. set up specific properties for each printer if you need.

Follow this procedure for each printer:

1.) Select Utility--> Connectivity--> Service. Add the Standard Print.



Figure 3-14 Add the Printer

2.) Highlight Standard Print in the Service list. Select the printer from the Printer pull-down Properties menu. For the UP-D897 printer, select "Portrait" as orientation. Type the printer name in the Name field. This name is used on the Button screen. After you select the printer from the Printer pull-down Properties menu again, it turns white. Press **Save**.

TCP/IP Device	Service D	taflow Button	Removable Media	Miscellaneous
Destination Device MyCo	omputer =			
Standard Print	-		Properties	
Station of Print	AUG	Printer	Sony UP-D897	
Service		Rows	1.	
Copy to Dataflow		Columns	1.	
HD Export Local Archive - Int HD		Orientation	Portrait	
Standard Print	Remove	Top Margin (mm)	0 *	
USB Drive H		Bottom Margin (mm)	0 -	
USD MUCK BUYE		Left Margin	0 .	
Propertie	15	Right Margin	0 -	
ame Standard Print	8			

Figure 3-15 Select the Printer

 Select *Button*. Select the appropriate print key (Print, Store...) from the Physical Print Buttons section. Select the printer from the MyComputer column and press >> to move it to the Printflow View column. Press *Save*.



Figure 3-16 Select Button
Section 3-3 - Receiving and Unpacking the Equipment

3-6-5-5 Digital Printer Instructions

Follow these steps to set up the paper size of the printer, take Sony UP-D897 as an example.

1.) Press Utility-->System-->Peripherals. Select the UP-D897 from the pull-down menu under Standard Printer Properties. Click *Properties*.



Figure 3-17 Properties

2.) Select **Properties** from Printer pull-down menu.

the Dataset See 1					
cument Native	Salut	Owner	Pages	504	Submitted

Figure 3-18 Properties

3.) Click Printing Preferences at the bottom of Properties Window.



Figure 3-19 Printing Preferences

- 4.) Select Paper Size. Press Apply. Press OK.
- 5.) Press Save, then Exit.

3-6-6 Available Probes

See in specification in the LOGIQ V5/LOGIQ V3 User Reference Manual for Probes and intended use.

Probe Name	Material of Headshell	Area of Using	TYPE	Catalog Number	Part Number
4C-RS	NORYL	GENERAL PURPOSE	CONVEX	H4000SR	5451471
E8C-RS	VALOX	TRANSVAGINAL TRANSRECTAL	MICRO-CONVEX	H40402LN	5409293
3Sc-RS	VALOX	CARDIAC	SECTOR	H45041DL	47237516
L6-12-RS	VALOX	SMALL PARTS PERIPHERAL VASCULAR	LINEAR	H48062AC	5454332

Table 3-16 List of Probes on LOGIQ V5/LOGIQ V3

Section 3-7 Software/Option Configuration

Refer to the LOGIQ V5/LOGIQ V3 Basic User Manual, Chapter 16, Customizing Your System for information on configuring items like Hospital, Department, Language, Units (of measure), Date, Time and Date Format.

For information on configuring Software Options, Refer to the LOGIQ V5/LOGIQ V3 Basic User Manual, Chapter 16, Customizing Your System.

For information on configuring DICOM Connectivity, Refer to the LOGIQ V5/LOGIQ V3 Basic User Manual, Chapter 16, Customizing Your System.

Section 3-8 Connectivity Installation Worksheet

Site System Information	
Site: Dept: LOGIQ SN: Type:	Floor: Comments: Room:
CONTACT INFORMATION Name Title	Phone E-Mail Address
TCP/IP Settings Name - AE Title: IP Settings IP Address: Subnet Mask: Default Gateway:	Remote Archive Setup Remote Archive IP: Remote Archive Name:
Device Type Manufacturer Name 1	IP Address Port AE Title

Section 3-9 Loading Base Image Software

Refer to

Section 8-3 "Loading Base Image Software" on page 8-4.

Section 3-10Software Version check out

3-10-1 Functional Check-out

- 1.) Power on LOGIQ V5/LOGIQ V3 scanner and wait until system booting to main screen.
- 2.) Press Utility key on control panel.
- 3.) Choose the About button on the right.



Figure 3-20 About and Software version

4.) Check whether "Software version" is the right version for use.

Section 3-11 Paperwork

NOTE: During and after installation, the documentation (i.e. User Manuals, Installation Manuals...) for the peripheral units must be kept as part of the original system documentation. This will ensure that all relevant safety and user information is available during the operation and service of the complete system.

3-11-1 Product Locator Installation

NOTE: The Product Locator Installation Card shown may not be same as the provided Product Locator card.

Maiing Address	GE Medical Systems Product Locator File P.O. Box 414 Milwaukee, WI 53201-0	0414	Ge Pro 28. 78.	neral Electr oduct Locat 3 Route de 530 Buc, FR	ic CGR or Adm I la Miniere ANCE	DSE/SM	Yoko GEM 4-7-1 Hino-	gawa Me SA Servio 27 Asahi -shi Toky	edical Systems Ltd. ce Administration igaoka o 191, JAPAN
DESCRIPTION		FDA	MO	CEL			REV	SERIAL	
SYSTEM UD.		┥		OCP	BS	ORD			EMLOYEE NO.
				OISTRICT	ROOM				DATE (MO - DA - YR)
IN IOT				CUSTOMER N	0.				J
INSI	ALLAIIU	Ν		DESTINATION NAME AND ADDRESS				12.5 - 20 -	
				8					
				3					
46-303268 R	ev 5			0.					ZIP CODE

Figure 3-21 Product Locator Installation Card

3-11-2 User Manual(s)

User Check that the correct User Manual(s) for the system and software revision, is included with the installation. Specific language versions of the User Manual may also be available. Check with your GE Sales Representative for availability.

Chapter 4 Functional Checks

Section 4-1 Overview

4-1-1 Purpose for Chapter 4

This chapter provides procedures for quickly checking major functions of the LOGIQ V5/LOGIQ V3 console, diagnostics by using the built-in service software, and power supply.

Table 4-1 Contents in Chapter 4

Section	Description	Page Number
4-1	Overview	4-1
4-2	Required Equipment	4-1
4-3	General Procedure	4-2
4-4	Software Configuration Checks	4-32
4-5	Peripheral Checks	4-32

Section 4-2 Required Equipment

To perform these tests, you'll need any of the sector, linear, or convex transducers.

(normally you should check all the transducers used on the system)

Section 4-3 General Procedure

CAUTION SYSTEM REQUIRES ALL COVERS

Operate this unit only when all board covers and frame panels are securely in place. The covers are required for safe operation, good system performance and cooling purposes.

4-3-1 Power On/Boot Up

After connect the system to the electrical supply, the power is applied to the scanner. When the Control panel *Power On/Off* key is pressed once, the System starts.

4-3-1-1 Scanner Power On

When power is applied to the scanner, power is distributed to the Cooling Unit, Control Panel, LCD, Peripherals and the Back-end Processor.

4-3-1-2 Turn on the system

Press the *Power On/Off* switch at the front of the system once.



Figure 4-1 Power On/Off Switch

When the *Power On/Off* switch on the Control Panel is pressed once, the Back-end Processor starts and the software code is distributed to initiate the scanner.

No status messages are displayed during this process.

4-3-2 Power Off/ Shutdown

NOTE: After turning off a system, wait at least ten seconds before turning it on again. The system may not be able to boot if power is recycled too quickly.

4-3-2-1 Back-end Processor Power Down

To power down the system:

- 1.) Press the *Power On/Off* switch at the front of the system once.
- 2.) The System-Exit window is displayed.

	Logon Information	_
Syr	stem Administrator is logged on as ADM	
ogon Time	2014-05-08 - 16:16	
S	oftware Remote Upgrade Information	=
Softwa	are Download Service connection failed	

Figure 4-2 System-Exit Window

- 3.) Using the Set key, select Shutdown.
- 4.) The shutdown process takes a few seconds and the power off sequence is complete when the power status LED is turned blue.
- 5.) Disconnect the probes. Clean or disinfect all probes as necessary. Store them in their shipping cases to avoid damage.

4-3-2-2 Scanner Shutdown

Disconnect the Mains Power Cable is necessary. For example: Relocating the scanner.

CAUTION DO NOT unplug and/or transport the unit until after the power off sequence has been completed. Failure to do so may result in corrupted patient files.

4-3-2-3 Check System Date and Time

A warning message "Please check the system date and time are correct" appears on the screen when the system is powered on. This warning message appears for the possible reasons:

- The system is not boot up for over 14 days.
- The system time has been changed by 24 hours earlier than the current system time of last boot-up.
- The BIOS time is changed by 24 hours earlier than the current system by resetting BIOS time, replacing BIOS module or changing BIOS time.

This warning message is to remind the user to check the system date in case the system date and time is incorrect.



Figure 4-3 Check system date and time message

Move the cursor to **OK** and press **Cursor** key on the control panel to select **OK**. The system enters scanning mode.

Check the system date and time. If it is incorrect, follow below steps to reset the system date and time.

- 1.) Enter Utility -> System -> General -> Date/Time.
- 2.) Reset the system date and time.
- 3.) Select Apply and then select OK.
- 4.) Select Save.

4-3-3 Archiving and Loading Presets

NOTE: Always save presets before any software reload. This ensures the presets loaded after the software reload are as up–to–date as possible.

All user presets except changes to Summary, Anatomy, and Biometry pages, can be saved on an CD-R disk (or USB memory device) for reloading on the system.

NOTICE Presets should NOT be saved on the same CD-R disk (or USB memory device) as images. The Archive Menu lists the images but does NOT list the presets stored on a CD-R disk (or USB memory device).

4-3-3-1 Archiving Presets to an CD-R Disk (or USB memory device)

- 1.) Insert an empty (blank) CD-R disk into the DVD-RW.
- 2.) Access to the **Utility** Menu, and select **System**. The Backup sheet will be shown on the LCD display.



Figure 4-4 Backup Sheet

- 3.) Select the item to back up either from Resource Files.
- 4.) Enter backup destination or browse through the disk to locate the destination.
- 5.) Select Backup now. The backup status for each item is displayed on the Result column.

4-3-3-2 Loading Presets from an CD-R disk (or USB memory device)

- 1.) Insert the CD-R disk with the archived Presets into the DVD-RW.
- 2.) Access to the Utility Menu, and select System. The Restore sheet will be shown on the LCD display.
- 3.) Select the item to restore either from Resource Files.
- 4.) Enter restore destination or browse through the disk to locate the destination.
- 5.) Select Restore. The restore status for each item is displayed on the Result column.

4-3-4 Adjusting the Display Monitor

Please refer to Section 6-2 "Monitor Adjustments" on page 6-2

4-3-5 System Features

4-3-5-1 Control Panel



Figure 4-5 Control Panel Tour

1.	Power On/Off	11. End Exam key	21. M/D Cursor key
2.	Rotary Button	12. Archive key	22. Scan Area key
3.	Page Up/Down keys	13. Scan Coach	23. Set /B Pause key
4.	TGC	14. Mode/Gain	24. B Steer/ Depth key
5.	AN Keyboard	15. Cursor key	25. AO and CHI key
6.	User Defined Keys	16. Clear key	26. Left/Right key
7.	Patient/Preset key	17. Comment key	27. Freeze key
8.	Probe key	18. Body Pattern	28. Print key
9.	User Preset key	19. Measure key	29. Store key
10.	Worksheet key	20. Ellipse/ Zoom key	

4-3-5-2 LOGIQ V5/LOGIQ V3 SoftMenu Key Tour

- 1.) Rotary key: Rotate to adjust the menu.
- 2.) Page Up/Down key: To turn the menu page up and down.



Figure 4-6 SoftMenu Key Tour

4-3-5-3 Monitor Display





- 1. Institution/Hospital Name, Date, Time, Operator Identification.
- 2. Patient Name, Patient Identification.
- 3. Power Output Readout.
- 4. Probe Identifier. Exam Preset.
- 5. Imaging Parameters by Mode.
- 6. Cine Gauge.
- 7. Active Images screen.
- 8. Delete Image.
- 9. Save As Menu.
- 10. Number of Images in Exam.
- 11. Page Indication.
- 12. Touchpad Functionality Status.
- Current date and time, Caps Lock: (lit when on), network connection indicator (PC=connected, PC with X=not connected), system messages display, InSite status, InSite controls. Image Preview

- 14. Image Preview.
- 15. Measurement Summary Window.
- 16. Worksheet/Direct Report.
- 17. Probe Orientation Marker.
- 18. Image.
- 19. Region of interest.
- 20. Gray/Color Bar.
- 21. Measurement Results Window.
- 22. Image Clipboard.
- 23. Measurement Calipers.
- 24. TGC.
- 25. Depth Scale.
- 26. Focal Zone Indicator.
- 27. Body Pattern.

4-3-6 B Mode Checks

4-3-6-1 Preparations

- 1.) Connect one of the probes listed in 3-6-6 "Available Probes" on page 3-24, in Chapter 3 Installation to the System probe connector.
- 2.) Turn ON the scanner (if it isn't turned on already).



Figure 4-8 Controls available in B Mode



Figure 4-9 B Mode Screen Picture Example

Table 4-2B Mode Controls

		Description/Benefit
Control	Possible Bioeffect	
Depth	Yes	Depth controls the distance over which the B-Mode images anatomy. To visualize deeper structures, increase the depth. If there is a large part of the display which is unused at the bottom, decrease the depth.
Gain	No	B-Mode Gain increases or decreases the amount of echo information displayed in an image. It may have the effect of brightening or darkening the image if sufficient echo information is generated.
Focus	Yes	Increases the number of focal zones, moves the focal zone(s) and change the zone width so that you can tighten up the beam for a specific area. A graphic caret corresponding to the focal zone position(s) appears on the right edge of the image.
Auto Optimize	No	Auto Optimize (Auto) lets you optimize the image based upon a the actual B Mode image data (Auto Tissue Optimize, ATO). The preset levels (Low, Medium, and High) allow you to pick a preference for the contrast enhancement in the resulting image. Low does the least amount of contrast enhancement, high does the most. Auto is available in single or multi image, on live, frozen or CINE images (in B-Mode only), and while in zoom and in Spectral Doppler. Auto in PW Doppler Mode(ASO) optimizes the spectral data. Auto adjusts the Velocity Scale/PRF (live imaging only), baseline shift, dynamic range, and invert (if preset). Upon deactivation, the spectrum is still optimized.
Mode Cursor	No	Displays the M/D-Mode cursor on the B-Mode image.
CrossXBeam	Yes	CrossXBeam is the process of combining three or more frames from different steering angles into a single frame. CrossXBeam is available on Convex and Linear probes. CrossXBeam combines multiple co-planar images from different view angles into a single image at real-time frame rates, using bi-cubic interpolation.
Coded Harmonic Imaging (CHI)	Yes	Enhances image resolution and improved small parts imaging.
Frequency	Yes	Multi Frequency mode lets you downshift to the probe's next lower frequency or shift up to a higher frequency.
Steer	Yes	You can slant the B-Mode or Color Flow linear image left or right to get more information without moving the probe. The angle steer function only applies to linear probes.
Virtual Convex	Yes	On Linear and Sector probes, Virtual Convex provides a larger field of view in the far field.
TGC	No	TGC amplifies returning signals to correct for the attenuation caused by tissues at increasing depths. TGC slide pots are spaced proportionately to the depth. The area each pot amplifies varies as well. A TGC curve may appear on the display (if preset), matching the controls that you have set (except during zoom). You can choose to deactivate the TGC curve on the image.
Width	Yes	You can widen or narrow the size of the sector angle to maximize the image's region of interest (ROI).

		Description/Benefit
Control	Possible Bioeffect	
Tilt	Yes	You can steer the sector angle to get more information without moving the probe while in B-Mode, M-Mode, Doppler Mode, and Color Flow Mode.
Reverse	No	Flips the image 180 degrees left/right.
Dynamic Range	No	Dynamic Range controls how echo intensities are converted to shades of gray, thereby increasing the adjustable range of contrast.
Line Density	Yes	Optimizes B-Mode frame rate or spatial resolution for the best possible image.
Мар	No	The system supplies B, M, and Doppler Mode system maps.
Frame Average	No	Temporal filter that averages frames together, thereby using more pixels to make up one image. This has the effect of presenting a smoother, softer image.
Colorize	No	Colorize is the colorization of a conventional B-Mode image or Doppler Spectrum to enhance the user's ability to discern B, M, and Doppler Mode intensity valuations. Colorize is NOT a Doppler Mode. NOTE: You can colorize realtime or CINE images or Timeline CINE. Colorizes the gray scale image to enhance the eye's discrimination capability. Spectrum Colorize colorizes the spectrum as a function of power using the inverse of the Colorize map for the signal intensity in each Doppler line. Colorize enhances the visibility of the spectrum's characteristics and enhances your ability to identify spectral broadening and the edge contours of the spectrum used to define the peak frequency/velocity. The colorize bar displays while Colorize is activated.
Edge Enhance	No	Edge Enhance brings out subtle tissue differences and boundaries by enhancing the gray scale differences corresponding to the edges of structures.
Rotation	No	You can flip the image up/down. CAUTION: When reading an rotated image, be careful to observe the probe orientation to avoid possible confusion over scan direction or left/right image reversal.
Rejection	No	Selects a level below which echoes will not be amplified (an echo must have a certain minimum amplitude before it will be processed).
Suppression	No	Suppresses the noise in the image.

Table 4-2B Mode Controls

		Description/Benefit
Control	Possible Bioeffect	
SRI-HD	No	SRI-HD (Speckle Reduction Imaging High Definition) is an adaptive algorithm to reduce the unwanted effects of speckle in the ultrasound image. Image speckle usually appears as a grainy texture in otherwise uniform areas of tissue. Its appearance is related to image system characteristics, rather than tissue characteristics, so that changes in system settings, such as probe type, frequency, depth, and others, can change the appearance of the speckle. Too much speckle can impair image quality and make it difficult to see the desired detail in the image. Likewise, too much filtering of speckle can mask or obscure desired image detail. Extra care must be taken to select the optimal SRI-HD level. SRI-HD is available in 2D imaging and may be used with any transducer or clinical application when image speckle appears to interfere with the desired image detail.
LOGIQ View (Option)	No	 LOGIQ View provides the ability to construct and view a static 2D image which is wider than the field of view of a given transducer. This feature allows viewing and measurements of anatomy that is larger than what would fit in a single image. Examples include scanning of vascular structures and connective tissues in the arms and legs. LOGIQ View constructs the extended image from individual image frames as the operator slides the transducer along the surface of the skin in the direction of the scan plane. The quality of the resulting image is somewhat user-dependent and requires some additional skill and practice to develop proper technique and become fully proficient. LOGIQ View is not available for the following: Multi Image, Timeline Modes, Color Flow Mode or PDI Mode.

Table 4-2B Mode Controls

4-3-7 M Mode Controls

4-3-7-1 Preparations

- 1.) Connect one of the probes listed in 3-6-6 "Available Probes" on page 3-24, in Chapter 3 Installation to the System probe connector.
- 2.) Turn ON the scanner (if it isn't turned on already).



Figure 4-10 Controls available in M Mode



Figure 4-11 M Mode Screen Picture Example Section 4-3 - General Procedure

4-3-7-2 M Mode Controls

Control	Possible Bioeffect	Description/Benefit
Sweep Speed	No	Changes the speed at which the timeline is swept. Available in M-Mode, Doppler Mode and M Color Flow Mode.
Anatomical M- Mode (option)	Yes	Anatomical M-Mode gives you the ability to manipulate the cursor at different angles and positions. The M-Mode display changes according to a motion of the M cursor.

Table 4-3M Mode Controls

4-3-8 Doppler Mode Checks

4-3-8-1 Preparations

- 1.) Connect one of the probes listed in 3-6-6 "Available Probes" on page 3-24, in Chapter 3 Installation to the System probe connector.
- 2.) Turn ON the scanner (if it isn't turned on already).



Figure 4-12 Controls available in Doppler Mode



Figure 4-13 Doppler Mode Screen Picture Example

4 - 16

4-3-8-2 Doppler Mode Controls

Table 4-4 Dopple	r Mode Controls
------------------	-----------------

Control	Possible Bioeffect	Description/Benefit
Auto Spectral Optimize [ASO] (Auto)	Yes	Auto in Doppler Mode optimizes the spectral data. Auto adjusts the Velocity Scale/PRF (on live images only), baseline shift, Dynamic Range and invert (if preset). The benefit of Auto can be found in reduced optimization time and a more consistent and accurate optimization process.
Update	Yes	Toggles between simultaneous and update presentation while viewing the timeline.
Doppler sample volume gate position	Yes	Moves the sample volume gate on the B-Mode's Doppler Mode cursor. The gate is positioned over a specific position within the vessel.
Doppler sample volume length	Yes	Sizes the sample volume gate.
Scale (Velocity Scale)	Yes	Adjusts the velocity scale to accommodate faster/ slower blood flow velocities. Velocity scale determines pulse repetition frequency. If the sample volume gate range exceeds single gate Scale capability, the system automatically switches to high PRF mode. Multiple gates appear, and HPRF is indicated on the display.
Angle Correct	No	Estimates the flow velocity in a direction at an angle to the Doppler vector by computing the angle between the Doppler vector and the flow to be measured. <i>NOTE: When the Doppler Mode Cursor and angle</i> <i>correct indicator are aligned (the angle is O), you</i> <i>cannot see the angle correct indicator.</i>
Quick Angle	No	Quickly adjusts the angle by 60 degrees.
Wall Filter	No	Insulates the Doppler signal from excessive noise caused from vessel movement.
Baseline	No	Adjusts the baseline to accommodate faster or slower blood flows to eliminate aliasing.
Mode Cursor	No	Displays the Doppler Mode cursor on the B-Mode image.
Steer and Fine Steer	Yes	You can slant the ROI of the Color Flow linear image left or right to get more information without moving the probe. The angle steer function only applies to linear probes.
Volume	No	Controls audio output.
Invert	No	Vertically inverts the spectral trace without affecting the baseline position.
Compression	No	Dynamic range controls how echo intensities are converted to shades of gray, thereby increasing the range of contrast you can adjust.
Trace Method	No	Traces the average mean and peak velocities in realtime or frozen images.
Cycles to Average	No	The average value over a number of cycles (from 1-5).
Trace Sensitivity	No	Adjust the trace to follow the waveform for signal strength.
Trace Direction	No	Specifies trace direction.

Chapter 4 - Functional Checks

Table 4-4	Doppler Mode Controls
-----------	-----------------------

Control	Possible Bioeffect	Description/Benefit
Display Format	No	Changes the horizontal/vertical layout between B-Mode and M-Mode, or timeline only.
Modify Auto Calcs	No	Activates the menu to select which calculations are automatically calculated.
Auto Calcs	No	Activates the calculation automatically which you select in the Modify Auto Calculation when the system is in a state of freeze or live.
Simultaneous (Duplex/Triplex)	Yes	When you select Duplex or Triplex, everything is live. For example, both B-Mode and PW/CW Doppler Modes are active if Duplex is selected. When Triplex is selected, B-Mode, PW/CW Doppler Mode, and CF Doppler Modes are active. If Duplex/Triplex is not selected, use M/D Cursor or Update to toggle between modes.

4-3-9 Color Flow Mode Checks

NOTE: Color Flow Mode only support by LOGIQ V5.

4-3-9-1 Preparations

- 1.) Connect one of the probes listed in 3-6-6 "Available Probes" on page 3-24, in Chapter 3 Installation to the System probe connector.
- 2.) Turn ON the scanner (if it isn't turned on already).



Figure 4-14 Controls available in Color Flow Mode



Chapter 4 - Functional Checks

4-3-9-2 Color Flow Mode Controls

Table 4-5 Color Flow Mode Controls

Control	Possible Bioeffect	Description/Benefit	
Flow Selection	No	In the Lower Extremity Vein (LEV) and Abdominal applications, you can quickly select the flow state via a shortcut on the Color Flow Mode menu.	
Gain	No	Gain amplifies the overall strength of echoes processed in the Color Flow window or spectral Doppler timeline.	
Scale (Velocity Scale)	Yes	Increases/decreases the Scale on the color bar.	
Wall Filter	No	Filters out low flow velocity signals. It helps get rid of motion artifacts caused from breathing and other patient motion.	
Size/Position	Yes	Adjust size and position of the color window.	
Invert (Color Invert)	No	Lets you view blood flow from a different perspective, e.g., red away (negative velocities) and blue toward (positive velocities). You can invert a real-time or frozen image. <i>NOTE: Invert reverses the color map, NOT the color PRF.</i>	
Baseline	No	Changes the Color Flow or Doppler spectrum baseline to accommodate higher velocity blood flow. Minimizes aliasing by displaying a greater range of forward flow with respect to reverse flow, or vice versa. Baseline adjusts the alias point. The default baseline is at the midpoint of the color display and at the midpoint of the color bar reference display.	
Angle Steer	Yes	You can slant the ROI of the Color Flow linear image left or right to get more information without moving the probe. The Angle Steer function only applies to linear probes.	
Accumulation	No	Accumulation enhances the flow in an image. Available in Color Flow, PDI, and B Flow.	
Color Flow Line Density	Yes	Optimizes the Color Flow frame rate or spatial resolution for the best possible color image.	
Мар	No	Allows you to select a specific color map. After you have made your selection, the color bar displays the resultant map.	
Map Compress	No	Change the gradation of color map.	
Threshold	No	Threshold assigns the gray scale level at which color information stops.	
Frame Average	No	Averages color frames.	
Transparency Map	No	Brings out the tissue behind the color map.	
Spatial Filter	No	Smooths out the color, makes it look less pixely.	
Flash Suppression	No	Activates/deactivates Flash Suppression, a motion artifact elimination process.	
Packet Size	Yes	Controls the number of samples gathered for a single color flow vector.	
Sample Volume	Yes	Adjusts the size of the color flow doppler transmit wave (or pulse) and size (or length). Lower setting gives better flow resolution and a higher setting increases sensitivity.	

Control	Possible Bioeffect	Description/Benefit
CF/PDI Auto Sample Volume	Yes	Set the default value at Utility -> Imaging -> CF Mode.
CF/PDI Focus Depth	Yes	
CF/PDI Frequency	Yes	
CF/PDI Auto Frequency	Yes	
CF/PDI Center Depth	Yes	
PDI	Yes	Power Doppler Imaging (PDI) is a color flow mapping technique used to map the strength of the Doppler signal coming from the tissue rather than the frequency shift of the signal. Using this technique, the ultrasound system plots color flow based on the number of reflectors that are moving, regardless of their velocity. PDI does not map velocity, therefore it is not subject to aliasing.
TVI (Option)	Yes	Tissue Velocity Imaging (TVI) calculates and color-codes the velocities in tissue. The tissue velocity information is acquired by sampling of tissue Doppler velocity values at discrete points. The information is stored in a combined format with gray scale imaging during one or several cardiac cycles with high temporal resolution.
TVD	Yes	TVD: Tissue Velocity Doppler: basing on TVI mode, activate a sample volume of PW ventricular wall to get the spectral information of the sample section.

Table 4-5 Color Flow Mode Controls

4-3-10 Basic Measurements

NOTE: The following instructions assume that you first scan the patient and then press **Freeze**.

4-3-10-1 Distance and Tissue Depth Measurements

- 1.) Press Measure once, an active caliper displays.
- 2.) To position the active caliper at the start point (distance) or the most anterior point (tissue depth), move the **Trackball**.
- 3.) To fix the start point, press **Set/ B Pause**. The system fixes the first caliper and displays a second active caliper.
- 4.) To position the second active caliper at the end point (distance) or the most posterior point (tissue depth), move the **Trackball**.
- 5.) To complete the measurement, press **Set**. The system displays the distance or tissue depth value in the measurement results window.

Before you complete a measurement:

To toggle between active calipers, rotate Cursor Select button.

To erase the second caliper and the current data measured and start the measurement again, press **Clear**.

- NOTE: To rotate through and activate previously fixed calipers, rotate **Cursor Select** button.
- NOTE: After you complete the measurement, to erase all data that has been measured to this point, but not data entered onto worksheets, press **Clear**.

4-3-10-2 Circumference/Area (Ellipse) Measurement

- 1.) Press **Measure** once; an active caliper displays.
- 2.) To position the active caliper, move the Trackball.
- 3.) To fix the start point, press **Set/ B Pause**. The system fixes the first caliper and displays a second active caliper.
- 4.) To position the second caliper, move the **Trackball**.
- 5.) Rotate the **Ellipse** button; an ellipse with an initial circle shape appears.
- NOTE: Be careful not to press the Ellipse control as this activates the Zoom.
 - 6.) To position the ellipse and to size the measured axes (move the calipers), move the Trackball.
 - 7.) To increase/ decrease the size, Rotate the Ellipse button.
 - 8.) To toggle between active calipers, *rotate Cursor Select button*.
 - 9.) To complete the measurement, press *Set/ B Pause*. The system displays the circumference and area in the measurement results window.

Before you complete a measurement:

- To erase the ellipse and the current data measured, press *Clear* once. The original caliper is displayed to restart the measurement.
- To exit the measurement function without completing the measurement, press *Clear* a second time.

4-3-10-3 Worksheets

Measurement/Calculation worksheets are available to display and edit measurements and calculations. There are generic worksheets as well as Application specific worksheets. The worksheets are selected from the Measurement Touch Panel.

4 - 22

4-3-10-4 Report Pages

Measurements/Calculations that are included on the worksheet can also be displayed on Report Pages. Report Pages can be customized to meet the appropriate needs of the user.

4-3-11 **Probe/Connectors Usage**

4-3-11-1 Connecting a probe

- 1.) Place the probe's carrying case on a stable surface and open the case.
- 2.) Carefully remove the probe and unwrap the probe cable.
- 3.) DO NOT allow the probe head to hang free. Impact to the probe head could result in irreparable damage.
- 4.) Align the connector with the probe port and carefully push into place.
- 5.) Lock the probe.
- 6.) Carefully position the probe cord so it is free to move and is not resting on the floor.

4-3-11-2 Activating the probe

The probe activates in the currently-selected operating mode. The probe's default settings for the mode and selected exam are used automatically.

4-3-11-3 Deactivating the probe

- 1.) Press the *Freeze* key.
- 2.) Gently wipe the excess gel from the face of the probe. (Refer to the User Guide for complete probe cleaning instructions.)
- 3.) Carefully slide the probe around the right side of the keyboard, toward the probe holder. Ensure that the probe is placed gently in the probe holder.

4-3-11-4 Disconnecting the probe

Probes can be disconnected at any time. However, the probe should not be selected as the active probe.

- 1.) Unlock the probe.
- 2.) Pull the probe and connector straight out of the probe port.
- 3.) Carefully slide the probe and connector away from the probe port and around the right side of the keyboard.
- 4.) Ensure the cable is free.
- 5.) Be sure that the probe head is clean before placing the probe in its storage box.

WARNING Take the following precautions with the probe cables: Do not bend, be sure to keep probe cables free from the wheels.

4-3-12 Using Cine

4-3-12-1 Activating CINE

Press **Freeze**, then roll the **Trackball** to activate CINE. To start CINE Loop playback, rotate **Run/Stop** button. To stop CINE Loop playback, rotate **Run/Stop** button.

4-3-12-2 Quickly Move to Start/End Frame

Rotate *First* button to move to the first CINE frame; rotate *Last button* to move to the last CINE frame.

4-3-12-3 Start Frame/End Frame

Rotate the *Start Frame* button to move to the beginning of the CINE Loop. Rotate the *Start Frame* clockwise to move forward through the CINE Loop. Rotate the *Start Frame* anticlockwise to move backward through the CINE Loop.

Rotate the *End Frame* button to move to the end of the CINE Loop. Rotate the *End Frame* button clockwise to move forward through the CINE Loop. Rotate the *End Frame* anticlockwise to move backward through the CINE Loop.

4-3-12-4 Adjusting the CINE Loop Playback Speed

Rotate the Loop Speed button to increase/decrease the CINE Loop playback speed.

4-3-12-5 Moving through a CINE Loop Frame By Frame

Rotate the *Frame by Frame* button to move through CINE memory one frame at a time.

4-3-13 Backup and Restore Database, Preset Configurations and Images

4-3-13-1 Formatting Media

- 1.) To format the backup media, enter Utility-> Connectivity->Removable Media.
- 2.) Select the media type from the drop down menu.
- 3.) Enter the label for the media as shown in Figure 4-16 on page 4-25. It is best to use all capital letters with no spaces or punctuation marks. Select **Format**.

Removable Media USB Driver H:	Verify		
AYAN	Format		
and the second s	Quic	k Format 💌	
Propert	ies		
Capacity 7367.0	MB		
Free space 7149.8	MB		
Formatted Yes			
Database Present Yes			
DICOMDIR Present No			
Finalized (CD Only)			
Write Protected No			
CD/DVD Type USB S	torage		
COUDUR Frank Time			

Figure 4-16 Format and Verify Media

4.) The system displays a pop-up menu, as shown in Figure 4-18 on page 4-26, select **OK** to continue.



Figure 4-17 Format Warning Pop-up Window

5.) If desired, verify that the format was successful by returning to **Utility-> Connectivity->Removable Media** and selecting **Verify** as shown in Figure 4-16 on page 4-25.

4-3-13-2 Backup System Presets and Configurations

- NOTE: Always backup any preset configurations before a software reload. This ensures that if the presets need to be reloaded, after the software update, they will be the same ones the customer was using prior to service.
 - 1.) Insert a formatted media into the drive.
 - 2.) Enter enter Utility-> System-> Backup/Restore.
- NOTE: If you are not logged in as GE Service or with administrator privileges, the Operator Login window is displayed. Log on with administrator privileges.
 - 3.) In the Backup list, select **Patient Archive**, **Report Archive**, **User Defined Configuration** and **Service**.
 - 4.) In the Media field, select CD/DVD (or USB memory device).
 - 5.) Select Backup, as shown in Figure 4-18 on page 4-26.

The system performs the backup. As it proceeds, status information is displayed on the Backup/Restore screen.



Figure 4-18 Backup/Restore Menu
4-3-13-3 Restore System Presets and Configurations

- **CAUTION** The restore procedure **overwrites** the existing database on the local hard drive. Make sure to insert the correct CD (or USB memory device).
 - 1.) Insert the Backup/Restore CD/DVD (or USB memory device) into the drive.
 - 2.) Enter enter Utility-> System-> Backup/Restore.
 - NOTE: If you are not logged in with administrator privileges, the Operator Login window is displayed. Log on with administrator privileges.
 - 3.) In the Restore list, select **Patient Archive**, **Report Archive**, **User Defined Configuration** and **Service**.
 - 4.) In the Media field, select the Backup/Restore CD/DVD (or USB memory device).
 - 5.) Select Restore.

The system performs the restore. As it proceeds, status information is displayed on the Backup/Restore screen.

General	System Imaging	System Measure	Backup/ Restore	Peripherals	User Configurable Key	About
		Backup			Restore	
User Define For Report 1	Patient Archive Report Archive d Configuration Service templates, use U	 Finished (Finished (Finished (Finished (Finished (Jtility/Report/E 	DK Thu May 08 DK Thu May 08 DK Thu May 08 DK Thu May 08 xport	17:23:22 2014 17:23:49 2014 17:23:88 2014 17:25:02 2014	Patient Archive V Report Archive V er Defined Configuration V Service V Restore	
Backup					Detailed Restore of User D	efined
Media USB	Drive H 💌	Media			Imag Connectivity Co	ing Presets
		EZMove			Measurement Co	onfiguration 🗌
Move Files Media capac	older Than in M M ity for estimate	Days 7 👻 Iedia CD / DVD (MB) 650 💌]=	R	Comment/Body Patte port Templates (Same Software Vo Utility->Applicat	rn Libraries ersion Only) Fast Key tion Presets
		EZBackup			Custo	m Programs
Reminder Ena	Dialog Interval i ble Reminder D M	Days 1 -			Restore	An others

Figure 4-19 Backup/Restore Menu

4-3-13-4 Archiving Images

- 1.) Insert the archive media.
- 2.) To format the archive media, enter Utility-> Connectivity-> Removable Media.
- 3.) Format the CD. Verify the format if desired.
- 4.) Images will be moved from the hard drive by date. Therefore, the best way is to label media by date.
- NOTE: Images will be moved from the hard drive by date. Therefore, the best way to label media is by date. When images are moved to the archive media, they will be deleted from the system hard drive. However, the patient database (backed up earlier) maintains pointers to the location of the images on the archive media.

TCP/IP Device Se	ervice Dataflow	Button
Removable Media	Verify	
Label	Format	
	Quick Format	
Properti	ies	
Capacity 7367.0	MB	
Free space 7149.8	MB	
Database Present Yes		
DICOMDIR Present No		
Finalized (CD Only)		
Write Protected No		
CD/DVD Type USB S	torage	
CD/DVD Storage Type		

Figure 4-20 Format Media Screen

- 5.) Enter Utility-> System-> Backup/Restore,
- 6.) Select "Move File Older Than in Days".



Figure 4-21 EZBackup/Move

Section 4-3 - General Procedure

4-3-13-4 Archiving Images (cont'd)

7.) Go to Archive menu and select EZBackup/EZMove, the EZBackup/EZMove wizard starts.

	Bearin Segu	Patient (D	a string	10.000	Clear .		Links Mar 24
9	Patrone at	Last Name	Paul Instan	Bellevin		I Last Public	COMPANY AND IN
Archive View	- 441212.464	100				20121110 11 22	3.52 MB
EL.	= ##1212-#58	111	1.1	Concernance of the		2012081317.92	2.05 MB
1.86	993912-833	Ob-ates	14	10720101	184	2012/10/10 11:26	62.1 848
Active Incases	= 112					2912/06/28 68:37	Plantes
	201200100					2012/06/18 15:02	33.7 MB
and the second second	201200178		4		1.04	2012/08/17 08:52	24.8 658
Data Tinesilar	#16	Text.	test			2012/06/28 11:52	None
	= B1. 891413	44			14	2012/08/17 14:12	24.3 108
	#5_0916t3	44				2012/08/14 18:55	0.01 MB
	#5_091712	42			N.	2012/08/17 14:17	0.01 MB
	85 091713	41			- 14	2912/09/17 14:35	0.03 MB
#2Nather	85 001713	41				3912/98/17 14:37	None
	. R.R. 346743	AIL			- 14	TRADING'S 7 15 PE	LAT MR
Arrite	Terrar .		Archive - Int. HD		U.	pin.	
Antes	Random Rectific		Antine - M. HD		0	bin .	
Acrim	Reading				10	bia	
Review	Reading Sector Sector Sector Sector		Anitina - N. HD		1	bia	
Perine	Baladine Second Baladine Baladine Baladine Alimet (16),000				1	him	
Review	Baladhuar Sanadhuar Balanacht Brussen Althuar (162000					him	
Berne	Batalture Versitäri Missiona Missiona Alling Picker				1	Lin .	
Receive.	Database sector Base of Base of Base Arrise caldes		Anthre - 14, 160			in a	
Arrite	Database Section Secti					Lin .	
Arrite	Distribut United at United at United at United at United at		Anthree HL HD			Lin .	
Arrite	Brinner Sternen Desterner Alter Ficker		Anitive - Bit. HD			Rim.	
. Poster	Distriction University Conversity Officers Conversity Conversity		Archive - bit, HD			Lin .	
Arrity	Bindhine Sintenia Waters Bindenia Attis fundes		Andrew - MA, HD			Rim.	

Figure 4-22 Archive Screen

8.) Verify the information on the first page of EZBack/EZMove wizard, then select Next.

If you want to backup all of the exams in the range (even if the exam was previously backed up), check this option. If you uncheck this option, the system only backs up exams which have not yet been backed up.

Welcome to EZBack	kup Wizard 🖾
-unstitling	Welcome to the GE Ultrasound E2Backup wizard!
1333	It has been 4706 day(s) since last back up.
And in case of the local division of the loc	Currently there is no active exam running.
	Local Images
Summer of the local division of the local di	12 Backup magen older Barr Bosy
	Euff backup
	Destination drive: Removable CD Archive Please review backup options. Click Next to continue
	Back Next Cancel

Figure 4-23 EZBackup Wizard 1

4-3-13-4 Archiving Images (cont'd)

- 9.) It indicates the size of the data and the storage. Select **Next** to continue.
- NOTE: The calculation for the number of backup CD is only an estimate. Allow for one additional CD when performing an EZBack/EZMove.

torage Size Information	
Storage Size Information Please review the detail storage size in each se	ction and prepare blank discs
Size of Data :	
Putient Archive :	4 MD
Images to back up :	31.8 MD
Number of images larger than 565 Mil	
Total nize :	36.8 MB
New discs needed (approximated):	t (each disc capacity is 650 MB)
	Back Next Cancel

Figure 4-24 EZBackup Wizard 2

NOTE: This message "Please insert a blank media..." appears if you press Next without inserting the backup media. Insert the media and continue.



Figure 4-25 Insert Media Message

4-3-13-4 Archiving Images (cont'd)

10.) The status menu appears. When the backup/move has been complete, press Next.

Please insert disk whe	n prompted		8
System is backing up	data		
Scanning images Skip 44 non-exist Formating disc 20 Formating disc 20	for oversit ng image(121119_02 121119_02	te kninges s). 	N 19
Disc Serial Humber :	20121119	02	
Progress:			3
Disc status : For	mating dis	c 20121119_02_	
Total Image Illumber:	3	Total Image Size:	31.9 MB
Image Done:		Image Done Size:	0.0 MH

Figure 4-26 EZBackup Wizard 3

- NOTE: If you need to insert the next media, a message appears providing you with the media label. Label the media, then insert the next media and press OK.
 - 11.)When the backup is complete, the completion wizard page appears. Press **Finish**.

Completion of EZBac	kup Wizard 🔯
	flackup completed.
	Please store the following disc(s) in a safe a place:
9	
	Back Finish Cancel

Figure 4-27 EZBackup Completion Window

All databases, presets and images should now be saved to removable media.

Section 4-4 Software Configuration Checks

Table 1-6	Software	Configuration	Chacks
1 abie 4-0	Sollware	Configuration	CHECKS

Step	Task to do	Expected Result(s)
1.	Check Date and Time setting	Date and Time are correct
2.	Check that Location (Hospital Name) is correct	Location Name is correct
3.	Check Language settings	Desired Language is displayed
4.	Check assignment of Printer Keys	The default function for Store and Print Keys are Store (store image), Print (print). Store and Print Keys can also be assigned as desired by the customer
5.	Check that all of the customer's options are set up correct	All authorized functions are enabled

Section 4-5 Peripheral Checks

Check that peripherals work as described below:

Table 4-7	Peripheral Checks
-----------	-------------------

Step	Task to do	Expected Result(s)
1.	Press (FREEZE)	Stop image acquisition.
2.	Press (PRINT) on the Control Panel	The image displayed on the screen is printed on B&W printer.
3.	Connect with Foot Switch on USB port and press once.	To start image acquisition (the same function as (FREEZE) key).

This page was intentionally left blank.

Chapter 5 Components and Functions (Theory)

Section 5-1 Overview

5-1-1 Purpose of Chapter 5

This chapter explains LOGIQ V5/LOGIQ V3 system concepts, component arrangement, and subsystem function.

Section	Description	Page Number
Section 5-1	Overview	5-1
Section 5-2	Block Diagram	5-2
Section 5-3	Common Service Platform	5-3

Table 5-1 Contents in Chapter 5

Section 5-2 Block Diagram

5-2-1 System Diagram





Section 5-3 Common Service Platform

5-3-1 Introduction

The Service Platform contains a set of software modules that are common to all PC backend ultrasound and cardiology systems. The Common Service Platform will increase service productivity and reduce training and service costs.

This page was intentionally left blank.

Chapter 6 Service Adjustments

Section 6-1 Overview

6-1-1 Purpose of this chapter 6

This section describes how to test and adjust the scanner. These tests are optional. You may use them to check the system for errors.

Table 6-2Contents in chapter 6

Section	Description	Page Number
Section 6-1	Overview	6-1
Section 6-2	Monitor Adjustments	6-2

Section 6-2 Monitor Adjustments

6-2-1 Adjustments Procedures

To adjust the brightness: Press **Home** and **End** button on the keyboard.



Figure 6-2 Light Adjustment

Chapter 7 Diagnostics/Troubleshooting

Section 7-1 Overview

7-1-1 Purpose of Chapter 7

This section describes how to setup and run the tools and software that help maintain image quality and system operation. Very basic host, system and board level diagnostics are run whenever power is applied. Some Service Tools may be run at the application level. However most software tests are required.

Section	Description	Page Number
7-1	Overview	7-1
7-2	Gathering Trouble Data	7-2
7-3	Screen Captures	7-4
7-4	Common Diagnostics	7-7
7-5	Network Configuration	7-8

-

 Table 7-3
 Contents in Chapter 7

Section 7-2 Gathering Trouble Data

7-2-1 Overview

There may be a time when it would be advantageous to capture trouble images and system data (logs) for acquisition to be sent back to the manufacturer for analysis. There are different options to acquire this data that would give different results.

7-2-2 Collect Vital System Information

The following information is necessary in order to properly analyze data or images being reported as a malfunction or being returned to the manufacturer:

- Product Name = LOGIQ V5/LOGIQ V3

From the *Utility>System>General>About* screen:

Applications Software

- Software Version
- Software Part Number

System Image Software

- Image Revision
- Image Part Number

7-2-3 Collect a Trouble Image with Logs

If the system should malfunction, press the Alt-D keys simultaneously. This will collect a screen capture of the image monitor, system presets and the following logs:

- Keyboard Shadow Log
- Error Logs
- Crash Log
- Power Supply
- Temperature
- NOTE: Power Supply and Temperature logs are not currently being updated by the LOGIQ V5/LOGIQ V3.

This Alt-D function is available at all times.

_	Export stored in	operta	
Description of is Address the full 11 Date and time 21 Sequence of 4 31 in this repeats Address the full 41 imaging mode 51 Bledis brand. 61 Dave seconds	sua i velig i el eccurance veent leading to izone de 7 veent estimating to izone h probe, presetrappication speet, capacity, type (eg, CD-R, DVD+RW ry image capitale, stine head, 40 multi volu	(etc.) anne longe	
System lucku) (application has been instarted after pro In the date and times when the problem (blem) scrutred.	
Contract Water			
Destination.	CD / DVD Recordable (G:)	Store	

Figure 7-1ALT-D Dialog Box

When Alt-D is pressed, a menu box appears that allows for:

- A place to enter a description of the problem
- A choice to store to a pre-formatted CD-R, RD (Removable Disk) or to the *Export* directory D: drive.

The subsequent file is compressed and time stamped. The screen capture is a bitmap which eliminates the possibility of artifacts from compression.

Section 7-3 Screen Captures

There may be times when the operator or field engineer will want to capture a presentation on the screen. This is accomplished by first saving the image(s) to the clipboard using a Print Key.

7-3-1 Check and Record the Store Key Function

Check the function of the Store Key in the event that the customer may have made some custom settings.

- 1.) Press *Utility* on the keyboard.
- 2.) Select *Connectivity* from the Utilities Menu.
- 3.) Select the *Button* tab on the **Connectivity** screen.
- 4.) In the *Physical Print Buttons* field, select Store.

The Connectivity/Buttons Screen will be displayed as shown in Figure 7-2 on page 7-4.

TCP/IP Device	Service	Dataflow	Estter/	Removabl	e Media	Miscellaneous
Physical Print B Print3 Print Store	uttons	MyComp Cop Star HD USE	uter ny to Dataflow ndard Print Export 3 Quick Save	er B My	Printflow \ ntflow View Computer 2 Copy to Dataf	low
Format Dicom (*.	dem)	L				
mane Frames Seconda						

Figure 7-2 Define Store Key Operation

If Store key is not set to Whole Screen, as shown in Figure 7-2 on page 7-4, proceed to step 5 to record the customer's customized settings.

- 5.) In the Destinations section, record the service that is displayed.
- 6.) In the *Physical Print Buttons* section, record the parameters related to the service.

7-3-2 Setting the Store Key to Screen Capture

If the Store Key is not set to screen capture:

- 1.) While on the Connect screen, with the Buttons tab displayed, go to the Destinations list.
- 2.) From the list select *Copy To Dataflow*. Press [>>] to add the selection to the *Printflow View* section.
- 3.) Ensure that the *Physical Print Buttons* section for capture Area is set to Whole Screen, secondary Capture and No Image Compression.
- 4.) The Store Key should now be set up for whole screen capture, sending the screens to the image buffer (clipboard).

7-3-3 Capturing a Screen

The following is a generic process to capture any screen from the scanner:

- 1.) Navigate to and display the image/screen to be captured.
- 2.) Press **Store**. This will place a snapshot of the screen on the "clipboard" displayed at the bottom of the scan image display.



Figure 7-3 Select Image to Capture

- 3.) Press **Freeze** to unfreeze the image to view the image screen and the snapshots displayed on the bottom.
- 4.) Highlight the snapshot to be stored.
- 5.) Select the save as icon on the right side of the image screen.

7-3-3Capturing a Screen (cont'd)

6.) A Save dialog box will be opened. Choose *d*:*export folder* as the archive location to save the image on the hard disk or CD.

		SAVE AS		
Save in ar	hive US	B Drive H: exp	ort	
Folder name		File name	Image01	
Store	• Image	only		
	Second	dary capture		
	(Children and Chil		1000	
Compression	Jpeg			
Compression Quality	Jpeg 100			Save
Compression Quality Save as type	Jpeg 100 Jpeg (Jp	9)		Save
Compression Quality Save as type	Jpeg 100 Jpeg (*)p	9)		Save
Compression Quality Save as type Delete Files For 1	Jpeg 100 Jpcg (Ljp	g)	Transl	Save Cancel

Figure 7-4 Save Dialog Box

NOTICE After capture the snapshot of the screen to the "clipboard" and save it to the hard disk or other media, it is not full screen image on the hard disk or media.

7-3-4 Reset the Store Key to Customer's Functionality

If the customer had programmed the Store Key to a function other than screen capture, restore that functionality recorded in 7-3-2 "Setting the Store Key to Screen Capture" on page 7-4. Refer to Figure 7-2.

- 1.) Click *Utility* on the keyboard.
- 2.) Select *Connectivity* from the Utilities Menu.
- 3.) Select the *Buttons* tab on the Connectivity screen.
- 4.) In the *Physical Print Button* field, select Store.
- 5.) In the Destinations list, select the service(s) recorded in step 5, Section 7-3-1.
- 6.) In the *Physical Print Buttons* section, select the parameters related to the service recorded in step 6, Section 7-3-1.

Section 7-4 Common Diagnostics

7-4-1 Utilities

Provides two selections:

7-4-1-1 Disruptive Mode

Allows you to enable or disable disruptive mode troubleshooting.

7-4-1-2 System Shutdown

Allows for system shutdown from the diagnostic menu. Select to *Restart System* or *Shutdown System*. Also, select to retain Disruptive Mode or Not.

After submitting to restart or shutdown a confirmation screen gives one last chance to confirm or cancel the request.

Section 7-5 Network Configuration

7-5-1 Network Configuration

- 1.) Connect system with network.
- 2.) Enter Utility-> Connectivity-> TCP/IP, in IP settings window, check Enable DHCP, and select the proper network speed in Network Speed.

TCP/IP Dev	vice Service Dat	aflow Button	Removable
Computer Name	LOGIQF		
	IP settings		0
Enable DHCP	•		
IP-Address			
Subnet Mask			
Default Gateway	1 1		
Network Speed:	Auto Detect		
Restart the syste	Auto Detect 10Mbps/Half Duplex 10Mbps/Full Duplex 100Mbps/Half Duplex 100Mbps/Full Duplex	aved from this page!	
	100Mbps/Full Duplex 1000Mbps/Auto-negotiate		

Figure 7-5 Enable DHCP

NOTE: If user wants to setup static IP address, uncheck **Enable DHCP** option, input static address in **IP-Address box**, **Subnet Mask** and **Default Gateway** box. In **Network Speed**, choose the proper speed available.

		IP settin	32		
Enable	DHCP -				Ŧ
IP-Ad	mess 3.35.8	8.3			
Subnet	Mask 255.25	i5.255.0			
efault Gat	eway 3.35.8	8.250			
Network S	peed: Auto I	Detect			
Restart the	system to a	ctivate any ch	anges saved f	rom this page!	
ce attaint this	a) menti to a	AMAAA SADDINA	anyes sarres i	Contraction product	

Figure 7-6 Input static address

7-5-1 Network Configuration (cont'd)

3.) Select **Save**, and a popup window displays. Select **OK** to restart the system and activate the changes.



Figure 7-7 System Restart inquiry dialog

4.) After the system restarts, the network icon at the left bottom of screen turns green.



Figure 7-8 Network icon

This page was intentionally left blank.

Chapter 8 Replacement Procedures

Section 8-1 Overview

8-1-1 Purpose of Chapter 8

This chapter describes replacement procedures for the following modules and subsystems.

Table 8-1Contents in Chapter 8

Section	Description	Page Number
8-1	Overview	8-1
8-2	DISASSEMBLY/RE-ASSEMBLY	8-2
8-2-1	Warning and Caution	8-2
8-2-2	Returning/Shipping for repairs	8-2
8-3	Loading Base Image Software	8-4

-

Section 8-2 DISASSEMBLY/RE-ASSEMBLY

8-2-1 Warning and Caution

- WARNING ONLY QUALIFIED SERVICE PERSONNEL SHOULD REMOVE ANY COVERS OR PANELS. ELECTRICAL HAZARDS EXISTS AT SEVERAL POINTS INSIDE. BECOME THOROUGHLY FAMILIAR WITH ALL HAZARDOUS VOLTAGES AND HIGH CURRENT LEVELS TO AVOID ACCIDENTAL CONTACT
- CAUTION Do not wear the ESD wrist strap when you remove a part of power supply unit. Turn OFF power and unplug the power cord before removing a part of power supply unit. However be sure to turn off power and wear the strap before you remove a circuit boards.

WARNING DO NOT SERVICE OR DISASSEMBLE PARTS UNDER FRU UNIT LEVEL AT ANY CIRCUMSTANCES.

8-2-2 Returning/Shipping for repairs

Equipment being returned must be clean and free of blood and other infectious substances.

GE policy states that body fluids must be properly removed from any part or equipment prior to shipment. GE employees, as well as customers, are responsible for ensuring that parts/equipment have been properly decontaminated prior to shipment. Under no circumstance should a part or equipment with visible body fluids be taken or shipped from a clinic or site (for example, body coils or an ultrasound probe). The purpose of the regulation is to protect employees in the transportation industry, as well as the people who will receive or open this package.

NOTE: The US Department of Transportation (DOT) has ruled that "items that were saturated and/or dripping with human blood that are now caked with dried blood; or which were used or intended for use in patient care" are "regulated medical waste" for transportation purposes and must be transported as a hazardous material.

If the LOGIQ V5/LOGIQ V3 needs to be sent for repair, ensure that any patient information is erased from the Harddisk/Storage Device. In case that any patient information is still residing on the LOGIQ V5/ LOGIQ V3, GE will contact the customer and request for urgent collection of that patient information. GE will keep this patient information in a secure environment for a maximum period of 1 month. All patient information will be permanently deleted at that point.

If PHI (Patient Healthcare Information) data needs to be sent to GE employees for service purposes, GE will ascertain agreement from the customer. The patient information shall only be transferred by approved service processes, tools and devices restricting access, protecting or encrypting data where required, and providing traceability in the form of paper or electronic documents at each stage of the procedure while maintaining compliance with cross-border restrictions of patient information transfers.

8-2-3 Footrest

8-2-3-1 Tools

Common phillips screwdrivers

8-2-3-2 Preparations

• Shut Down the System and disconnect the power cord.

8-2-3-3 Removal procedure

1.) Unscrew 4 screws at the bottom side, and remove the footrest, refer to Figure 8-9 on page 8-3.



Figure 8-9 Footrest Cover Disassembly

-

8-2-3-4 Mounting procedure

Install the new parts in the reverse order of removal.

Section 8-3 Loading Base Image Software

- NOTE: While it is believed to be unnecessary, It would not hurt to disconnect the system from the network and remove all transducers.
- NOTE: Please ensure power cable is connected during system upgrade!
 - 1.) Insert the USB disk labeled "System & Application Software" in the USB port of the system.
 - 2.) Properly turn off the scanner by momentarily pressing the *Power On/Off* Switch. Select "Shut Down" from the System Exit menu.
 - 3.) If the system will not shutdown normally, hold down the *Power On/Off* Switch until the light turns from green to amber.
 - 4.) Turn on the system. System will detect the USB automatically.
 - 5.) Press any key to continue when below message display as shown below.

DRENING - DAENING - WAENING - WAENING - WAENING - WAENING	
THIS PROCEDURE MAY RESULT IN COMPLETE PATIENT BATA LOSS IT NOT USED Correctly? Please read the options below carefully before proceeding.	
This process is NOT REVERSIBLE and should NOT be stopped once started? DO NOT power off the system until the process has completed. It will take less than 28 minutes to load the drive. IF this process IS stopped for some reason, you WILL have to run it again to completion or else the system will not work.	
If yow want to proceed with this process press the "Enter" key to continue with option selection.	P)
Remove the USB memory stick and Press "CTRL+Alt+Bel" now to exit and power cycle your system to restart it without overwrifing your disk drive's current contents.	
Press any key to continue	

Figure 8-10 Upgrade message

Section 8-3 Loading Base Image Software (cont'd)

6.) Select one of the options for loading the system. Select choice 1] to load the complete disk.



Figure 8-11 Selection for loading the system

- WARNING While the software install procedure is designed to preserve data, you should select choice [2] to format disk C only.
 - 7.) Press any key to continue when below message display as shown below.



Figure 8-12 Upgrade continue message

Section 8-3 Loading Base Image Software (cont'd)

8.) System will be loaded as shown in the screen below.



Figure 8-13 System Load

9.) System updating finished, refer to Figure 8-14 on page 8-6.



Figure 8-14 System upgrade complete

10.)Remove the USB stick, then press power key to restart system.

Chapter 9 Renewal Parts

Section 9-1 Overview

9-1-1 Purpose of Chapter 9

This chapter gives you an overview of Renewal Parts for LOGIQ V5/LOGIQ V3.

Table 9-1Contents in Chapter 9

Section	Description	Page Number
9-1	Overview	9-1
9-2	List of Abbreviations	9-1
9-3	Renewal Parts Lists	9-2
9-4	Operator Console Assy	9-3
9-5	CRU Parts List	9-4
9-5	Manuals	9-5
9-7	Peripheral	9-7

Section 9-2 List of Abbreviations

- Assy Assembly
- Ctrl Control
- FRU 1 Replacement part available in part hub
- FRU 2 Replacement part available from the manufacturer (lead time involved)
- Int Internal
- I/O Input/Output
- LCD Liquid Crystal Display
- MON Monitor
- PAT. Patient
- PC Personal Computer (Back End Processor)

Section 9-3 Renewal Parts Lists

9-3-1 Equipment Models Covered in this Chapter

Table 9-2 ACDC Power Pack & ACDC Power Cable list

Part Name	Part Number	Description	Quantity	FRU
AC Power Cord	5177123-2	Power cord Europe Class	1	1
AC Power Cord	5176304-2	Power cord China Class	1	1
AC Power Cord	5176773-2	Power cord India Class	1	1
AC Power Cord	5177195-2	Power cord Argentina Class	1	1
AC Power Cord	5176907-2	Power cord UK Class	1	1
AC Power Cord	5177153-2	Power cord Denmark Class	1	1
AC Power Cord	5177154-2	Power cord Switzerland Class	1	1
AC Power Cord	5177187-3	Power cord Australia Class	1	1
AC Power Cord	5177146-2	Power cord USA Class	1	1
AC Power Cord	5176753-2	Power cord Israel Class	1	1
AC Power Cord	5177126-2	Power cord Japan Class	1	1
AC Power Cord	5400868-2	Power cord Brazil Class	1	1

Section 9-4 Operator Console Assy



Figure 9-1Operator Console Assy

Section 9-5 CRU Parts List

A CRU is a FRU part that can be replaced by the Customer and Customer comfirms functional check at site.

Part Name	Part Number	Description	Quantity	FRU
4C-RS Probe	5451471	4C-RS Convex Probe	1	1
E8C-RS probe	5409293	E8C-RS convex probe	1	1
3Sc-RS probe	47237516	3Sc-RS sector probe	1	1
L6-12-RS probe	5454332	L6-12-RS Linear probe	1	1
Power cord Europe Class	5177123-2	Power cord Europe Class	1	1
Power cord China Class	5176304-2	Power cord China Class	1	1
Power cord India Class	5176773-2	Power cord India Class	1	1
Power cord Argentina Class	5177195-2	Power cord Argentina Class	1	1
Power cord UK Class	5176907-2	Power cord UK Class	1	1
Power cord Denmark Class	5177153-2	Power cord Denmark Class	1	1
Power cord Switzerland Class	5177154-2	Power cord Switzerland Class	1	1
Power cord Australia Class	5177187-3	Power cord Australia Class	1	1
Power cord USA Class	5177146-2	Power cord USA Class	1	1
Power cord Israel Class	5176753-2	Power cord Israel Class	1	1
Power cord Japan Class	5177126-2	Power cord Japan Class	1	1
Power cord Brazil Class	5400868-2	Power cord Brazil Class	1	1

Table 9-3 CRU Parts List

9 - 4

Section 9-6 Manuals

Table 9-4

LOGIQ V Series Manuals

ltem	Part Number	Description	Qty	FRU		
7000	5496012-100	LOGIQ V5/LOGIQ V3 Basic Service Manual	1	N		
System User Manuals						
7001	5475853-100	LOGIQ V5/LOGIQ V3 Basic User Manual, English	1	N		
7002	5475843-100	LOGIQ V5/LOGIQ V3 User Guide, English	1	N		
7003	5475843-101	LOGIQ V5/LOGIQ V3 User Guide, French	1	N		
7004	5475843-106	LOGIQ V5/LOGIQ V3 User Guide, Spanish	1	N		
7005	5475843-108	LOGIQ V5/LOGIQ V3 User Guide, German	1	N		
7006	5475843-111	LOGIQ V5/LOGIQ V3 User Guide, Italian	1	N		
7007	5475843-121	LOGIQ V5/LOGIQ V3 User Guide, Dutch	1	N		
7008	5475843-127	LOGIQ V5/LOGIQ V3 User Guide, Brazilian Portuguese	1	N		
7009	5475843-129	LOGIQ V5/LOGIQ V3 User Guide, Estonian	1	N		
7010	5475843-131	LOGIQ V5/LOGIQ V3 User Guide, Slovenian	1	N		
7011	5475843-140	LOGIQ V5/LOGIQ V3 User Guide, Japanese	1	N		
7012	5475843-142	LOGIQ V5/LOGIQ V3 User Guide, Swedish	1	N		
7013	5475843-144	LOGIQ V5/LOGIQ V3 User Guide, Korean	1	N		
7014	5475843-145	LOGIQ V5/LOGIQ V3 User Guide, Russian	1	N		
7015	5475843-150	LOGIQ V5/LOGIQ V3 User Guide, Polish	1	Ν		
7016	5475843-151	LOGIQ V5/LOGIQ V3 User Guide, Greek	1	Ν		
7017	5475843-153	LOGIQ V5/LOGIQ V3 User Guide, Hungarian	1	Ν		
7018	5475843-154	LOGIQ V5/LOGIQ V3 User Guide, Slovakian	1	Ν		
7019	5475843-155	LOGIQ V5/LOGIQ V3 User Guide, Czech	1	Ν		
7020	5475843-159	LOGIQ V5/LOGIQ V3 User Guide, Turkish	1	N		
7021	5475843-160	LOGIQ V5/LOGIQ V3 User Guide, Danish	1	N		
7022	5475843-161	LOGIQ V5/LOGIQ V3 User Guide, Norwegian	1	N		
7023	5475843-162	LOGIQ V5/LOGIQ V3 User Guide, Finnish	1	Ν		
7024	5475843-165	LOGIQ V5/LOGIQ V3 User Guide, Bulgarian	1	Ν		
7025	5475843-167	LOGIQ V5/LOGIQ V3 User Guide, Romanian	1	N		
7026	5475843-168	LOGIQ V5/LOGIQ V3 User Guide, Croatian	1	N		
7027	5475843-174	LOGIQ V5/LOGIQ V3 User Guide, Lithuanian	1	Ν		
7028	5475843-175	LOGIQ V5/LOGIQ V3 User Guide, Latvian	1	N		
7029	5475843-176	LOGIQ V5/LOGIQ V3 User Guide, Serbian	1	N		
7030	5475843-177	LOGIQ V5/LOGIQ V3 User Guide, European Protuguese	1	N		
7031	5475843-181	LOGIQ V5/LOGIQ V3 User Guide, Indonesian	1	N		

Chapter 9 - Renewal Parts

Table 9-4LOGIQ V Series Manuals

ltem	Part Number	Description	Qty	FRU
7032	5498971-141	LOGIQ V3 User Guide, Chinese	1	Ν
7033	5498972-141	LOGIQ V5 User Guide, Chinese	1	Ν
Section 9-7 Peripheral

Table	9-5
-------	-----

5 LOGIQ V Model Designations

ltem	Part Name	Part Number	Qty	FRU		
	Printers					
8000	Sony UP-D897 Chinese kit	5151262	1	N		
8000A	Sony UP-D897 USA kit	5151259	1	N		
8000B	Sony UP-D897 European kit	5151261	1	N		
8000C	Sony UP-D897 Japanese kit	5151263	1	N		
8001	Sony UP-D25MD USA kit	5398062	1	N		
8001A	Sony UP-D25MD European kit	5398063	1	N		
8001B	Sony UP-D25MD Japanese kit	5398064	1	N		
8001C	Sony UP-D25MD Chinese kit	5398061	1	N		
8002	Sony UP-D711MD with Paper kit	5494719	1	N		
8002A	Sony UP-D711MD Paper 1 roll	5494718	1	N		
8002B	Sony UP-D711MD Printer Paper 10 rolls	5543548	1	N		
8003	HP Officejet 100 Chinese kit	5426594	1	N		
8003A	HP Officejet 100 European kit	5426595	1	N		
8003A	HP Officejet 100 Japanese kit	5426596	1	N		
8003B	HP Officejet 100 USA kit	5426597	1	N		
8004	Sony UP-D898MD USA kit	5151259-2	1	N		
8004A	Sony UP-D898MD European kit	5151261-2	1	N		
8004B	Sony UP-D898MD China kit	5151262-2	1	N		
8004C	Sony UP-D898MD Japan kit	5151263-2	1	N		
8004D	Sony UP-D898MD Brazil kit	5495509-2	1	N		
8005	HP Officejet Pro 8100	NA	1	N		
8006	Sony UP-D898MD Printer shelf option kit	5599283	1	N		
8007	Sony UP-D711MD Printer shelf option kit	5484791	1	N		
	Video	Converter				
8008	Video Converter Module	5534825	1	N		
	D\	/D-RW				
8009	LITEON eUAU108	5454614-2	1	N		
	For	otswitch				
8010	MKF 2-MED GP26 (IPx8)	5151236	1	N		
8011	FSU-1000 (IPx8)	5338419	1	N		
	US	B Stick				
8012	SanDisk CRUZER 4G	5395754	1	N		

Chapter 9 - Renewal Parts

Table 9-5 LOGIQ V Model Designations

Item	Part Name	Part Number	Qty	FRU
8013	1TB mobile USB HDD	5434317-3	1	N
	Bic	ppsy Kit		
8014	E8C-RS biopsy kit	E8385NA	1	N
8015	E8C-RS reusable biopsy kit	2398164	1	N
8016	4C-RS biopsy kit	5160703	1	N
8017	L6-12-RS biopsy kit	5176499	1	N
8018	3Sc-RS biopsy kit	5329137	1	N
	System and Appl	ication Software USB		•
8019	LOGIQ V Series R1.0.0 System and Application Software USB	5501727	1	Y
8020	LOGIQ V Series R1.0.1 System and Application Software USB	5501727-2S	1	Y
8021	LOGIQ V Series R1.0.2 System and Application Software USB	5501727-3S	1	Y
8022	LOGIQ V Series R1.0.3 System and Application Software USB	5501727-4S	1	Y

Section 9-8Probes

ltem	Part Name	Part Number	Description	Quantity	FRU
9000	4C-RS	5451471	Probe (Center Frequency: 3.10MHz)	1	1
9001	3Sc-RS	47237516	Probe (Center Frequency: 2.75MHz)	1	1
9002	L6-12-RS	5454332	Probe (Center Frequency: 7.75MHz)	1	1
9003	E8C-RS	5409293	Probe (Center Frequency: 6.5MHz)	1	1

Table 9-6 Probes on LOGIQ V5/LOGIQV3

NOTICE All the spare parts should be disposed according to local laws.

This page was intentionally left blank.

Chapter 10 Care & Maintenance

Section 10-1 Overview

10-1-1 Periodic Maintenance Inspections

It has been determined by engineering that your system does not have any high wear components that fail with use, therefore no Periodic Maintenance Inspections are mandatory. Some Customers Quality Assurance Programs may require additional tasks and or inspections at a different frequency than listed in this manual.

10-1-2 Purpose of Chapter 10

This chapter describes **Care & Maintenance** on the scanner and peripherals. These procedures are intended to **maintain the quality** of the ultrasound **systems performance**. Read this chapter completely and familiarize yourself with the procedures before performing a task.

Section	Description	Page Number
10-1	Overview	10-1
10-2	Why do Maintenance	10-2
10-3	Maintenance Task Schedule	10-2
10-4	Tools Required	10-4
Section 10-5	When There's Too Much Leakage Current	10-15

Table 10-1Contents in Chapter 10

- **CAUTION** Practice good ESD prevention. Wear an anti–static strap when handling electronic parts and even when disconnecting/connecting cables.
- DANGER THERE ARE SEVERAL PLACES ON THE BACKPLANE, THE AC DISTRIBUTION, AND DC DISTRIBUTION THAT ARE DANGEROUS. BE SURE TO DISCONNECT THE SYSTEM POWER PLUG AND OPEN THE MAIN CIRCUIT BREAKER BEFORE YOU REMOVE ANY PARTS. BE CAUTIOUS WHENEVER POWER IS STILL ON AND COVERS ARE REMOVED.
- **CAUTION** Do not pull out or insert circuit boards while power is ON.
- CAUTION Do not operate this unit unless all board covers and frame panels are securely in place. System performance and cooling require this.

-

Section 10-2 Why do Maintenance

10-2-1 Keeping Records

It is good business practice that ultrasound facilities maintain records of quality checks and corrective maintenance. The Ultrasound Inspection Certificate (provided on page 10-16) provides the customer with documentation that the ultrasound scanner is maintained on a periodic basis.

A copy of the Ultrasound Periodic Maintenance Inspection Certificate should be kept in the same room or near the scanner.

10-2-2 Quality Assurance

In order to gain accreditation from organizations such as the American College of Radiology (USA), it is the customer's responsibility to have a quality assurance program in place for each scanner. The program must be directed by a medical physicists, the supervising radiologist/physician or appropriate designee.

Routine quality control testing must occur regularly. The same tests are performed during each period so that changes can be monitored over time and effective corrective action can be taken.

Testing results, corrective action and the effects of corrective action must be documented and maintained on the site.

Your GE service representative can help you with establishing, performing and maintaining records for a quality assurance program. Please contact us for coverage information and/or price for service.

Section 10-3 Maintenance Task Schedule

10-3-1 How often should care & maintenance tasks be performed?

The Care & Maintenance Task Schedule (provided on page 10-3) specifies how often your LOGIQ V5/ LOGIQ V3 should be serviced and outlines items requiring special attention.

NOTE: It is the customer's responsibility to ensure the LOGIQ V5/LOGIQ V3 care & maintenance is performed as scheduled in order to retain its high level of safety, dependability and performance.

Your GE Service Representative has an in-depth knowledge of your LOGIQ V5/LOGIQ V3 ultrasound scanning system and can best provide competent, efficient service. Please contact us for coverage information and/or price for service.

The service procedures and recommended intervals shown in the Care & Maintenance Task Schedule assumes that you use your LOGIQ V5/LOGIQ V3 for an average patient load (10-12 per day) and not use it as a primary mobile unit which is transported between diagnostic facilities.

NOTE: If conditions exist which exceed typical usage and patient load, then it is recommended to increase the maintenance frequencies.

Section 10-3 Maintenance Task Schedule (cont'd)

Table 10-2 Customer Care Schedule

Service at Indicated Time	Daily	Weekly	Monthly	Per Facilities QA Program	Notes
Clean Probe Holders	•				
Clean Air Filter		•			more frequently depending on your environment
Inspect AC Mains Cable			•		Mobile Unit Check Weekly
Inspect Cables and Connectors			•		
Clean Console			•		
Inspect Wheels, Casters, brakes and Swivel Locks			•		Mobile Unit Check Daily
Check Control Panel Movement			•		Mobile Unit Check Daily
Console Leakage Current Checks				•	also after corrective maintenance
Peripheral Leakage Current Checks				•	also after corrective maintenance
Surface Probe Leakage Current Checks				•	also after corrective maintenance
Endocavity Probe Leakage Current Checks				•	also after corrective maintenance
Transesphongeal Probe Leakage Current Checks				•	also after corrective maintenance
Surgical Probe Leakage Current Checks				•	also after corrective maintenance
Measurement Accuracy Checks				•	also after corrective maintenance
Functional Checks				•	also after corrective maintenance

-

NOTE: PMs are not mandatory, the table above is for reference only.

NOTE: May require specialized equipment to complete.

Section 10-4 Tools Required

10-4-1 Standard GE Tool Kit

The following is a description of the "Standard" GE tool kit in the USA. Not all tools are required for PMs.

Table 10-3 Overview of GE-1 Tool Kit Contents

Tool ID	Description	Tool ID	Description
9-45358	Pliers Retaining Ring	9-XL9971MM	Xcelite-hex Blade 1.27mm
9-4078	Scribe	9-XL9972MM	Xcelite-hex Blade 1.5mm
9-44572	Wrench Open End 3/8 - 7/16	9-XL9973MM	Xcelite-hex Blade 2 mm
9-44579	Wrench Open End 1/2 - 9/16	9-XL9974MM	Xcelite-hex Blade 2.5mm
9-44579	Wrench Open End 1/2 - 9/16	9-XL9975MM	Xcelite-hex Blade 3mm
9-45385	Pliers, Arc Joint 7 inch	9-XL9976MM	Xcelite-hex Blade 4mm
9-45378	Pliers, Slip Joint	9-XL9977MM	Xcelite-hex Blade 5mm
9-4518	Pliers, Long Nose, Miniature	9-XL991CM	Handle
9-4518	Pliers, Long Nose, Miniature	C2356E	Screw starter - Kedman Quick Wedge
9-44776	Ignition Wrench Set, 10 pc.	BLBO	Box - 18 Compartment
9-44601	Wrench, Adj., 4 inch	DWL4283T	Box - 5 Compartment
9-4151	Screwdriver, Blade, Stubby	9-41322	Pickup Tool, Claw type
9-41421	Screwdriver, Blade, Pocket clip	9-6757	6 pc Needle File Set
9-41594	Screwdriver, Blade 1/8 in. x 4 in.	9-9487	Utility Knife
9-41581	Screwdriver, Blade 3/16 in. x 4 in.	9-45341	Pliers Vice Grip 10 inch
9-39451	20' Steel Tape, locking Spring load	9-3001	Xacto Pen Knife
9-GH807	Ratchet, Offset, Slotted	9-HT62002	Solder Aid, Fork and Hook
68-412	Ratchet, Offset, Phillips	9-4099	Mirror, Round, Telescoping
9-GH130	Tapered Reamer	9-GH3001	Steel Rule Decimal 6 inch
9-41584	Screwdriver, slotted 1/4 in.X 6 in.	9-GH300ME	Steel Rule Metric 6 inch
9-4118	Screwdriver, Phillips #2, Stubby	9-XL9920	Xcelite-hex Blade.050 inch
9-41293	Screwdriver, Phillips #0	9-XL9921	Xcelite-hex Blade 1/16 inch
9-41294	Screwdriver, Phillips #1	9-XL9922	Xcelite-hex Blade 5/16 inch
9-41295	Screwdriver, Phillips #2	9-XL9923	Xcelite-hex Blade 3/32 inch
9-46677	Hex Keys, 20 pc., Metric	9-XL9924	Xcelite-hex Blade 1/8 inch
9-34701	1/4 in. Standard.Socket set (19 pc)	9-XL9925	Xcelite-hex Blade 5/32 inch
9-43499	1/2 inch Socket 1/4 inch drive	9-XL9926	Xcelite-hex Blade 3/16 inch
9-4355	Flex Spinner	9-XL99764	Xcelite-hex Blade 7/64
9-43523	Breaker	9-XL99964	Xcelite-hex Blade 9/64
9-43531	6 inch Ext.	9-XLM60	Mini-screwdriver kit

Table 10-3	Overview of GE-1 Tool Kit Contents	(cont'd))
			£

Tool ID	Description	Tool ID	Description
9-65283	Case 8.5 in. x 4.5 in. x 2 in. Deep	9-45072	Pliers 6 inch Diagonal
9-46696	Hex Keys	9-XL100X	Wire Stripper/Cutter 5 inch - 100X
9-39829	Torpedo Level, Magnetic	9-XL87CG	Pliers - very fine needle nose-87CG
9-38461	Hammer, Ball Peen, 4 oz	9-WEWDT-07	Weller-Soldering-Replacement Tip(1)
9-4280	Universal Joint 1/4 inch	9-WS175-E	Wiss - Surgical Scissors
9-WEW60P3	Weller - Soldering Iron, 3 wire	KH174	Hemostat 5 inch Straight
9-WECT5B6	Weller - Soldering Iron Tip	KH175	Hemostat 5 inch curved
9-WEWDP12	Weller - Desoldering Pump	9-Z9480121	Alignment tool (red)
93383	Flashlight Mini-Mag Lite (AAA Bat.)		
9-GH408	Tweezers		
21576	Brush - Bristle		
9-4516	Pliers 4 1/4 inch Diagonal		

Table 10-4 Overview of GE-2 Tool Kit Contents(cont'd)

GE-2 Sears Kit (#99034)			
Tool ID	Description	Tool ID	Description
9-45381	Pliers, Arc Joint 9 1/2 inch	9-44067	Socket 1 1/16 in. for 1/2 in. drive
9-45092	Pliers, Linesman 8 1/2 inch	9-42679	Socket 10MM Hex for 1/2 in. drive (2273333)
9-42882	Punch, Pin 3/32 inch	9-44262	Extension 10 inch for 1/2 in. drive (2273405)
9-42884	Punch, Pin 5/32 inch	9-4258	3/8 inch to 1/2 inch Adapter
9-42886	Punch, Pin 1/4 inch	9-34374	3/8 inch Metric Socket Set - 12 PT
9-42973	Cold Chisel 1/2 inch	9-44311	16mm Socket 12 pt.
9-GH77	Center Punch Automatic	9-33485	Metal Socket Tray
9-GH890	File Handle, Adj.	9-33484	Metal Socket Tray
9-31276	File, Round, Bastard 8 inch	9-33484	Metal Socket Tray
9-31277	File, Half Round, Bastard 8 inch	9-52068	Tap and Drill Set
9-31263	File, Flat Mill 8 inch	9-52722	#6 Тар
21045C	Close Quarter Saw	9-52723	#8 Тар
9-44604	Wrench, Adj 10 inch		High Speed Drill Set
9-41587	Screwdriver 5/16 inch x 8 inch		#36 Drill
9-41586	Screwdriver, Stubby 5/16 inch		#29 Drill
9-GH19512	Countersink 1/2 inch	9-44046	3/8 inch Socket Set
9-44741	12 PC Combination Wrench Set		

10-4-2 Special Tools, Supplies and Equipment

10-4-2-1 Specific Requirements for Care & Maintenance

Table 10-5 Overview of Requirements for Care & Maintenance

ΤοοΙ	Part Number	Comments
Digital Volt Meter (DVM)		
Leakage Current Ultrasound Kit	2113015	For 120V and 220V Units
Anti Static Kit	46–194427P231 46–194427P279 46–194427P369 46–194427P373 46–194427P370	Kit includes anti–static mat, wrist strap and cables for 200 to 240 V system 3M #2204 Large adjustable wrist strap 3M #2214 Small adjustable wrist strap 3M #3051 conductive ground cord
Anti Static Vacuum Cleaner	46–194427P278 46–194427P279	120V 230V
Air Filter		air intake
Safety Analyzer		The Safety Analyzer tool should be calibrated and compliant with AAMI/ESI 1993 or IEC 60601 or AS/NZS 3551
SVHS VCR Cassette	E7010GG E7010GF	60 minute 120 minute
SVHS VCR Head Cleaner		See VCR user manual for requirements
3.5" MOD MEDIA	E8381AA E8381AB	blank 128 M disk blank 230 M disk
5.25" MOD Media		
3.5" MOD Media Cleaner	2117811	cleans the diskettes
5.25" MOD Media Cleaner		cleans the diskettes
3.5" MOD Head Cleaner Kit	2148392	cleans the drive heads
5.25" MOD Head Cleaner Kit		cleans the drive heads
QIQ Phantom	E8370RB	RMI Grayscale Target Model 403GS
B/W Printer Cleaning Sheet		See printer user manual for requirements
Color Printer Cleaning Sheet		See printer user manual for requirements
Disposable Gloves		

10-4-3 Input Power

10-4-3-1 Mains Cable Inspection

Table 10-6 Mains Cable Inspection

Step	ltem	Description
1	Unplug Cord	Disconnect the mains cable from the wall and system.
2	Inspect	Inspect it and its connectors for damage of any kind.
3	Verify	Verify that the LINE, NEUTRAL and GROUND wires are properly attached to the terminals, and that no strands may cause a short circuit.
4	Verify	Inlet connector retainer is functional.

10-4-4 Cleaning

10-4-4-1 General Cleaning

Table 10-7 General Cleaning

Step	ltem	Description
1	Console	Use a fluid detergent in warm water on a soft, damp cloth to carefully wipe the entire system. Be careful not to get the cloth too wet so that moisture does not enter the console.
2	Probe Holder	Clean probe holders (they may need to be soaked to remove excess gel).

NOTE: For your convenience or of the air filter is too dirty, replacement filters are available. refer to Chapter 9 for the air filter replacement part number.

10-4-5 Physical Inspection

	Table 10-	8 Ph	vsical (Checks
--	-----------	------	----------	--------

Step	ltem	Description
1	Labeling	Verify that all system labeling is present and in readable condition. refer to the LOGIQ V5/LOGIQ V3 User Manual for details.
2	Scratches & Dents	Inspect the console for dents, scratches or cracks.
3	Wheels & Brakes	Check all wheels and casters for wear and verify operation of foot brake, to stop the unit from moving, and release mechanism. Check all caster locks and caster swivel locks for proper operation.
4	Cables & Connectors	Check all internal cable harnesses and connectors for wear and secure connector seating. Pay special attention to footswitch assembly and probe strain or bend reliefs.
5	Shielding & Covers	Check to ensure that all EMI shielding, internal covers, air flow panels and screws are in place. Missing covers and hardware could cause EMI/RFI problems while scanning.
6	External I/O	Check all connectors for damage and verify that the labeling is good.
7	Op Panel Lights	Check for proper operation of all operator panel and TGC lights.
8	Monitor Light	Check for proper operation of any monitor lights if available.
9	External Microphone	Check for proper operation of any external microphones by recording an audio test.

Section 10-4 - Tools Required

10-4-6 Outlet Test -Wiring Arrangement - USA & Canada

Test all outlets in the area for proper grounding and wiring arrangement by plugging in the neon outlet tester and noting the combination of lights that are illuminated. Any problems found should be reported to the hospital immediately and the receptacle should not be used.



Figure 10-1 Typical Outlet Tester

NOTE: No outlet tester can detect the condition where the Neutral (grounded supply) conductor and the Grounding (protective earth) conductor are reversed. If later tests indicate high leakage currents, this should be suspected as a possible cause and the outlet wiring should be visually inspected.

10-4-7 Grounding Continuity

▲ CAUTION Electric Shock Hazard. The patient must not be contacted to the equipment during this test.

Measure the resistance from the third pin of the attachment plug to the exposed metal parts of the case. The ground wire resistance should be less than 0.2 ohms. Reference the procedure in the IEC 601-1.1.



Figure 10-2 Ground Continuity Test

10-4-7-1 Meter Procedure

Follow these steps to test the ground wire resistance.

- 1.) Turn the LOGIQ V5/LOGIQ V3 unit OFF.
- 2.) Plug the unit into the meter, and the meter into the tested AC wall outlet.
- 3.) Plug the black chassis cable into the meter's "CHASSIS" connector and attach the black chassis cable clamp to an exposed metal part of the LOGIQ V5/LOGIQ V3 unit.
- 4.) Set the meter's "FUNCTION" switch to the RESISTANCE position.
- 5.) Set the meter's "POLARITY" switch to the OFF (center) position.
- 6.) Measure and record the ground wire resistance.

10-4-8 Chassis Leakage Current Test

10-4-8-1 Definition

This test measures the current that would flow in a grounded person who touched accessible metal parts of the bedside station if the ground wire should break. The test verifies the isolation of the power line from the chassis. The meter is connected from accessible metal parts of the case to ground. Measurements should be made with the unit On and Off, with the power line polarity Normal and Reversed. Record the highest reading.

CAUTION Electric Shock Hazard. When the meter's ground switch is OPEN, don't touch the unit!

CAUTION Equipment damage possibility. Never switch the Polarity and the status of Neutral when the unit is powered ON. Be sure to turn the unit power OFF before switching them using the POLARITY switch and/or the NEUTRAL switch. Otherwise, the unit may be damaged.

10-4-8-2 Generic Procedure

The test verifies the isolation of the power line from the chassis. The testing meter is connected from accessible metal parts of the case to ground. Measurements should be made with the unit ON and OFF, with the power line polarity Normal and Reversed. Record the highest reading of current.



Figure 10-3 Set Up for Chassis Source Leakage Current, IEC 601-1 Clause 19 - Continuos Leakage Currents and Patient, Auxiliary Currents

When using the Microguard or a similar test instrument, its power plug may be inserted into the wall outlet and the equipment under test is plugged into the receptacle on the panel of the meter. This places the meter in the grounding conductor and the current flowing from the case to ground will be indicated in any of the current ranges. The maximum allowable limit for chassis source leakage is shown in Table 10-12.

10-4-9 Isolated Patient Lead (Source) Leakage–Lead to Lead

Reference the procedure in the IEC 60601-1. Select and test each of the five ECG lead positions (except ALL) on the LEAD selector, testing each to the power condition combinations found in the table. Record the highest leakage current measured.

10-4-10 Isolated Patient Lead (Sink) Leakage-Isolation Test

reference the procedure in the IEC 60601-1. Select the ALL position on the lead selector. Depress the rocker switch to ISO TEST to test lead isolation.

AUTION Line voltage is applied to the ECG leads during this test. To avoid possible electric shock hazard, the system being tested must not be touched by patients, users or anyone while the ISO TEST switch is depressed.

NOTE: It is not necessary to test each lead individually or power condition combinations as required in previous tests.

10-4-10-1 Data Sheet for ECG Leakage Current

The test passes when all readings measure less than the value shown in the table below. Record all data on the PM Inspection Certificate.

Table 10-9 Maximum Allowance Limit for ECG Leakage Current

		Maximum Allowance Limit		
	AC Power Source	GROUND OPEN	GROUND CLOSED	
Patient Lead to Ground Leakage Current Test	115V	10uA	10uA	
and Patient Lead to Lead Leakage Current Test	220/240V	500uA	10uA	

Table 10-10 Maximum Allowance Limit for ECG Leakage Current

	AC Power Source	Maximum Allowance Limit
Patient Lead Isolation Current Test	115V	20uA
	220/240V	5mA

Table 10-11 Typical Data Sheet for ECG Leakage Current

500	Tester	Tester	Tester Lead Selector					
Power	Switch	Switch	RL	RA	LA	LL	С	
ON	NORM	CLOSED						
ON	REVERSE	CLOSED						
ON	NORM	OPEN						
ON	REVERSE	OPEN						
OFF	NORM	CLOSED						
OFF	REVERSE	CLOSED						
OFF	NORM	OPEN						
OFF	REVERSE	OPEN						

10-4-11 Probe Leakage Current Test

10-4-11-1 Definition

This test measures the current that would flow to ground from any of the probes through a patient who is being scanned and becomes grounded by touching some other grounded surface.

10-4-11-2 Generic Procedure

Measurements should be made with the ground open and closed, with power line polarity normal and reversed, and with the unit Off and On. For each combination, the probe must be active to find the worst case condition.



Figure 10-4 Set Up for Probe Leakage Current

NOTE: Each probe will have some amount of leakage current, dependent on its design. Small variations in probe leakage currents are normal from probe to probe. Other variations will result from differences in line voltage and test lead placement.

10-4-11-3 Meter Procedure Using Probe Adapter

Follow the Safety Analyzer tool instruction to test each transducer for leakage current.

The electrical Safety Analyzer tool should be calibrated and compliant with AMMI/ESI 1993 or IEC 60601 or AS/NZS 3551.

10-4-11-4 No Meter Probe Adapter Procedure

Follow the Safety Analyzer tool instruction to test each transducer for leakage current.

The electrical Safety Analyzer tool should be calibrated and compliant with AMMI/ESI 1993 or IEC 60601 or AS/NZS 3551.

10-4-11-5 Data Sheet for Transducer Source Leakage Current

The test passes when all readings measure less than the values. Record all data on the PM Inspection Certificate.

CAUTION Equipment damage possibility. Never switch the Polarity and the status of Neutral when the unit is powered ON. Be sure to turn the unit power OFF before switching them using the POLARITY switch and/or the NEUTRAL switch. Otherwise, the unit may be damaged

Table 10-12 Typical Data Sheet For Transducer Source Leakage Current

Transducer Tested:						
Tester Power Polarity Tester GROUND or Unit Power Switch NEUTRAL Switch						
ON	NORM	OPEN				
ON	NORM	CLOSED				
ON	REV	OPEN				
ON	REV	CLOSED				
OFF	NORM	OPEN				
OFF	NORM	CLOSED				
OFF	REV	OPEN				
OFF	REV	CLOSED				

Section 10-5 When There's Too Much Leakage Current...

CHASSIS FAILS

Check the ground on the power cord and plug for continuity. Ensure the ground is not broken, frayed, or intermittent. Replace any defective part.

Tighten all grounds. Ensure star washers are under all ground studs.

Inspect wiring for bad crimps, poor connections, or damage.

Test the wall outlet; verify it is grounded and is free of other wiring abnormalities. Notify the user or owner to correct any deviations. As a work around, check the other outlets to see if they could be used instead.

NOTE: No outlet tester can detect the condition where the white neutral wire and the green grounding wire are reversed. If later tests indicate high leakage currents, this should be suspected as a possible cause and the outlet wiring should be visually inspected.

PROBE FAILS

Test the probe in another connector to isolate if the fault lies with the probe or the scanner.

NOTE: Each probe will have some amount of leakage, dependent on its design. Small variations in probe leakage currents are normal from probe to probe. Other variations will result from differences in line voltage and test lead placement. The maximum allowable leakage current for body surface contact probe differs from inter-cavity probe. Be sure to enter the correct probe type in the appropriate space on the check list.

If excessive leakage current is slot dependent, inspect the system connector for bent pins, poor connections, and ground continuity.

If the problem remains with the probe, replace the probe.

PERIPHERAL FAILS

Tighten all grounds. Ensure star washers are under all ground studs.

Inspect wiring for bad crimps, poor connections, or damage.

STILL FAILS

If all else fails, begin isolation by removing the probes, external peripherals, then the on board ones, one at a time while monitoring the leakage current measurement.

NEW UNIT

If the leakage current measurement tests fail on a new unit and if situation can not be corrected, submit a Safety Failure Report to document the system problem. Remove unit from operation.

ECG FAILS

Inspect cables for damage or poor connections.

ULTRASOUND INSPECTION CERTIFICATE

Customer Name: System Type		System ID: Dispatch Number / Date Performed:		Warranty/Contract/HBS	
		Model Number:	Serial Number:	Manufacture Date:	
Probe 1:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 2:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 3:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 4:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 5:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 6:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 7:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 8:	Frequency:	Scan Format*:	Model Number:	Serial Number:	
Probe 9:	Frequency:	Scan Format*:	Model Number:	Serial Number:	

* Scan Format: Phased Array, Linear Array, Curved Array, Mechanical Array or Other

FUNCTIONAL CHECKS

PHYSICAL INSPECTION AND CLEANING

Functional Check (if applicable)	OK? or N/A	Physical Inspection and Cleaning (if applicable)	Inspect	Clean
B-Mode Function		Console		
Doppler Modes Function		Monitor		
CF-Mode Function		Touch Panel		
M-Mode Function		Air Filter		
Applicable Software Options		Probe Holders		
Applicable Hardware Options		External I/O		
Control Panel		Wheels, Brakes & Swivel Locks		
Monitor		Cables and Connectors		
Touch Panel		GE Approved Peripherals (VCR, CD-RW, MOD, Printers)		
Measurement Accuracy				
GE Approved Peripherals				

COMMENTS:

10 - 16

ELECTRICAL SAFETY

Electrical Test Performed	Max Value Allowed	Value Measured	OK?	Comments
Outlet (correct ground &wiring config.)				
System Ground Continuity				
Chassis Source Leakage Current - Probe				
Chassis Source Leakage Current - Caster				
Chassis Source Leakage Current - CRT				
Patient Lead Source Leakage (Lead to Ground)				
Patient Lead Source Leakage (Lead to Lead)				
Patient Lead Source Leakage (Isolation)				
Peripheral 1 Leakage Current				
Peripheral 1Ground Continuity				
Peripheral 2 Leakage Current				
Peripheral 2Ground Continuity				
Peripheral 3 Leakage Current				
Peripheral 3Ground Continuity				
		PROBES		
Probe Number (from previous page)	Max Value Allowed	Max Value Measured	OK?	Comments
Probe 1:				
Probe 2:				
Probe 3:				
Probe 4:				
Probe 5:				
Probe 6:				
Probe 7:				
Probe 8:				
Probe 9:				

Final Check. All system covers are in place. System scans with all probes as expected.

Section 10-5 - When There's Too Much Leakage Current...

This page was intentionally left blank.

INDEX

A,B

Abbreviations, 9-1 Archiving Images Move, 4-28 Backup Patient Database, 4-25 Preset Configurations, 4-25 Basic Measurements Functional Checks, 4-22 Boot Up, 3-11

С

CE Compliance, 1-15 Color Mode Overview, 4-16 Connectivity Worksheet, 3-25 Contact Information, 1-16 Control Panel, 4-7 Customer Assistance, 1-16

D

Dangerous Procedure Warnings, 1-12 DICOM Network Function, 2-11

Е

Electrical Safety, 1-11 Electrostatic Discharge Warning, 1-15 EMI, 1-15 ESD, 1-15

F

facility needs suggested floor plan scanner and EchoPAC in same room, 2-10 floor plan suggestion scanner and EchoPAC in same room, 2-10 Functional Checks, 4-1 Basic Measurements, 4-22 Control Panel, 4-7 Monitor Display, 4-9 Peripherals, 4-32 Probes/Connector Usage, 4-23

G

Gathering Trouble Data, 7-2 General Cleaning, 10-7

Н

Hazard Icons, 1-6 Human Safety, 1-10

L LOTO, 1-14

м

Mechanical Safety, 1-10 Models Covered, 1-2 Monitor Display Functioanl Checks, 4-9 Move Archiving Images, 4-28

Ρ

P4 Key Function, 7-4 Peripherals Functional Checks, 4-32 Power On, 3-11 Power-up Procedures Voltage Settings, 3-10 Probes/Connector Usage Functional Checks, 4-23 Product Icons, 1-7

R

Required Features, 2-7 Restore Patient Database, 4-27 Preset Configurations, 4-27

S

Safety Considerations, 1-10 Screen Captures, 7-4 System Manufacturer, 1-17

Т

Index

Touch Panel Functional Checks, 4-7 Trouble Image with Logs, 7-3 Troubleshooting Gathering Trouble Data, 7-2 Screen Captures, 7-4 Trouble Image with Logs, 7-3 Vital System Information, 7-2

v

Voltage Settings, 3-10

W

Warnings and Cautions, 1-10

© 2014, General Electric Company. GE-GEMedical Systems Ultrasound. 9900 Innovation Drive Wawautosa, Wisconsin 53226 USA

