Development and preliminary evaluation of the VA Seattle foot


INTRODUCTION

Most mobility aids for physically handicapped individuals seek to restore and improve function that primarily relates to basic lifestyle needs. This is an appropriate priority. With the lower limb amputee, this objective means stable, bipedal standing, and walking on unobstructed level surfaces. These elementary needs should be accomplished with comfort and with reasonable energy output (7, 19). Presently available lower limb prostheses effectively satisfy these needs in most instances. However, as the mobility demands of an individual with amputation expand, conventional prostheses in general perform poorly. This circumstance is most evident when the amputee attempts to run. Incremental increases in speed through fast walking, jogging, and running rapidly cause gait alterations in which, with increasing speed, the unilateral amputee spends less and less time and weight on the deficient limb, which results in the sound limb largely propelling the body through the gait cycle. This resulting high-energy consuming, uncomfortable, unstable, and unsightly gait pattern is thus generally avoided. Very few bilateral leg amputees are able to run. For these reasons most amputees do not walk rapidly or run, and many have never even attempted to do so (5, 6, 13, 14).

The ability to move quickly and especially to run is a basic need for most physical recreation. It is also important to physical and mental well being and as a defense against injuries such as falling and avoiding threatening environmental situations.

The Prosthetics Research Unit has investigated in depth over the past seven years the running capabilities of a number of types and levels of lower limb amputees. Research kinesiologists of the University of Washington collaborated with the Prosthetics Research Unit by conducting extensive investigation over a 5-year period of amputee running (9, 10).

It became evident that the state-of-the-art lower limb prostheses would have to be redesigned if real progress in amputee running was to be accomplished. No amount of muscle conditioning and training alone could be expected to accomplish major change in running performance. The prosthetic foot, which is the subject of this article, is an outgrowth of this amputee gait performance research (Figs. 1A and 1B).

DESIGN OF THE SEATTLE FOOT

The VA Seattle foot conceptually provides storage of potential energy and is converted to
kinetic energy throughout the weight-bearing phase of the gait cycle. This stored energy is progressively released as the foot continues through the toe-off phase to rebound and propel the body forward. In simple terms, weight deflects the keel through a predetermined range, then the keel “springs back” as weight is removed (Fig. 2).

Specifications were determined mathematically using information from the gait studies and transferring it to the bench testing of keel materials which could meet requirements (1–3, 8, 11, 12, 15–18).

The first foot was fabricated from leaves of fiberglass combined with a light metal (4). When the foot was tried on patients, their response was one of remarkable acceptance. Continued use on test subjects brought out the problem areas: weight, breakage, individual amputee preferences for specific performance needs. After a series of refinements it became evident that the most satisfactory keel design would be monolithic. That is, with fewer component parts, the production cost would be lower, there would be less maintenance, and the response would be more uniform.

The present monolithic keel is composed of the acetal homopolymer, Dupont's Delrin®. Its design form and physical performance are described in the engineering section of this article. Foaming was initially in the general shape of a foot and suitable for shoe fitting. This shape corresponded to the one ordinarily used for commercially available prosthetic feet. As the design progressed, we decided to prepare anatomical molds so that the foot would actually resemble a natural foot for those people who desired this type of cosmesis. A few amputees did not wish this natural appearance and accordingly were fitting with the standard and previously used blank-foot shape (Fig. 3).

These criteria were selected to guide development beyond the proof-of-concept fiberglass epoxy keels: 1) store and return energy (1–3/4 inch metatarsal deflection at 455 pounds vertical load); 2) natural feel and stability; 3) useful life of 3 years; 4) lightweight (1 pound target); 5) reduced production costs.

An initial review of available space constraints within the foot shape, the large deflections required to simulate normal-foot A-P plane motions, and available material properties led to the following development approach: 1) uniform stress monolithic cantilever spring keel; 2) modular and compatible with existing prosthetic components (standard single-bolt attachment accessible from lower surface); 3) natural appearance; 4) minimize part count; 5) ultimate production keel to be molded; 6) fatigue: 50,000 cycles representing sprint running (load = 2.8 × body weight) – 1,000,000 cycles representing jogging (load = 1.4 × body weight); 7) less than 0.06 inch permanent set at 3 × body weight; 8) increase damping in the spring.

Early structural analysis of moldable materials showed that obtaining a sufficiently soft spring to allow required deflection at the forefoot was difficult, especially considering the demanding fatigue life requirements. Preliminary designs and structural analyses were conducted on nearly
100 combinations of thermoset and thermoplastic matrices and reinforcements. A constant width, parabolic tapered-depth beam was picked as the basic section for isotropic materials. Straight taper was used for oriented fiber composite candidates. A general keel shape consisting of a hook posterior to the bolt centerline and then descending to the metatarsals was selected because it would move the apparent “center of rotation” of the cantilever spring deflection as close as possible to the natural ankle center. It was also felt that increasing the outer extreme tension fiber length would improve fatigue life of the highly stressed keel materials. Load deflection testing of ¼ inch thick sections of the keel (“toes”) was conducted early to guide design and analysis. Toe testing allowed quick evaluation of material and process combinations for which mechanical properties were not available. Over 50 of these representative sections were tested (Fig. 4).

When required load-deflection characteristics were obtained, a fatigue test was run on the candidate material toe section. The 2-cycle-per-second fatigue test device is shown in Figure 5. The test load was applied 23 degrees anterior to the shank centerline, simulating maximum dorsiflexion loads at heel-rise and toe-off.

Of the several materials and shapes fatigue tested the Delrin® 150 material (DuPont’s acetal homopolymer) came closest to meeting all the criteria. Many materials which were strong enough were too stiff or had insufficient damping to allow a natural feel for the amputee. One of the shortcomings of the epoxy-fiberglass keel was that walking amputees felt “hurried” by the too-quick release of the stored spring energy. The significantly higher damping of Delrin® has eliminated this problem. Three-dimensional keels were then fatigue tested on the same device. A design was finally derived which met the fatigue criteria, although permanent set at the end of the conservative “straight-through” fatigue cycles was larger than desired (0.2 inch actual versus an objective of 0.06 inch). Subsequent patient testing on the natural intermittent loading cycle has not yet uncovered any creep problems.

**FIT PROTOCOL**

Clinical testing helped reveal flaws that have been corrected. Major problems areas were:
1) anterior keel tip (“toe”) failures, which was corrected by pre-bending to a larger tip-up angle and thinning the section to reduce stress induced by the approximately 60 degrees toe-up angle demanded during toe-off; 2) failures at the attachment bolt clearance hole in the bottom of the keel, corrected by providing additional reinforcing pad-up to carry loads around the hole; and 3) keel “punch-through” the foam on soft surface, e.g., barefoot in dry sand, corrected by adding a Kevlar® fabric toe pad (Fig. 6).

Eight keel configurations are currently required to cover the adult population of male shoe sizes 7 through 11 and body weights from 130 to 240 pounds. Use of the fit protocol shown in
Figure 7 has resulted in high satisfaction for individuals with amputation.

Selection of a keel for an exceptionally high activity level might require the next stiffer keel to avoid breakage or to provide better feel (higher energy storage). Similarly, a next softer keel might be prescribed for an inactive amputee. We have noted cases where an inactive patient (sometimes a bilateral amputee) has been fitted with a softer keel than body weight would dictate, and then found his activity level increasing to the point where he was able to break the keel. Since the keel is highly stressed during use by active amputees, all VA Seattle foot units are subjected to the rigorous acceptance test procedure presented in Figure 8.

Extensive acceptance clinical testing and evaluation was performed using 36 subjects (34 below-knee and 2 above-knee amputees). The high level of satisfaction/acceptance is recorded in Figures 9A, B, and C.

The foot was then placed into further evaluation with 500 volunteer amputees in 44 designated Veterans Administration Medical Centers across the United States. The feet were fitted to both new and currently worn prostheses by the subjects’ prosthetists of choice as designated by the clinic prosthetic teams at participating stations. This study is being conducted by the Rehabilitation Research and Development Evaluation Unit under the direction of James B. Reswick, Sc.D., in collaboration with the Veter-
FIGURE 4
Some of the keel designs and materials evaluated.

FIGURE 5
Bench-testing of keel for fatigue.

The Administration Central Office’s Prosthetics and Sensory Aids Service, which is directed by Frederick Downs, Jr. The Evaluation Unit was recently established by Margaret J. Giannini, M.D., as part of the Rehabilitation Research and Development Service. Data are now being collected and analyzed. Early unofficial reports indicate that they closely match the information derived from the small study.

A further commercial evaluation program was initiated and now completed using several hundred feet provided to prosthetists nationwide. This study, too, will be the subject of a subsequent report.
PROSTHETIC CONSIDERATIONS

The VA Seattle foot was designed for use with
conventional lower limb prostheses and is inter-
changeable with existing components. To accom-
plish this feature, the foot is attached to the
prosthetic shank, endoskeletal or exoskeletal,
with a single rigid bolt. Interchangeability, dura-
bility, simplicity, and cost were all addressed.
Earlier designs, which involved a number of
components, have been refined to the present
monolithic keel. Minimizing components reduces
servicing and cost. The VA Seattle foot has only
three components: the keel, the external foam,
and the toe reinforcement pad.

Weight of the appliance is a critical factor.
Currently, prosthetic feet weigh between \( \frac{1}{2} \) and
1½ pounds. The VA Seattle foot weighs approxi-
mately 1 pound, varying a small amount depend-
ing on the size used.

PRESCRIPTION CRITERIA

While the VA Seattle foot was initially de-
signed for running, it can also be used for walking
and is not necessarily contraindicated for less-
active people. Gait studies have shown that
because the foot is flexible in the metatarsal area,
it does not limit the forward travel of the tibia as
it rotates over the foot, thus making the transi-
tion between foot-flat and toe-off smoother. By
combining the ability to increase push-off
through deflection of the keel, the foot improves
both walking and running. When walking on
uneven ground, the VA Seattle foot does not
provide as much forefoot flexibility in the medial/
lateral plane as in the Greissinger or SAFE foot.
Therefore, if the patient requires this motion,
another foot should be selected. Continuing re-
search is being directed to increase component
compensation in additional planes as required by
irregular surfaces, inclines, and steps. We desire
to incorporate additional force-motion character-
istics within the material rather than add or
modify components.

As stated earlier, the VA Seattle foot can be
used with success on athletic and less-active
amputees, including individuals with bilateral
amputation. The foot can also be used on all levels

ALIGNMENT

Optimal performance of the VA Seattle foot is
generally more difficult to achieve when it is
attached to an existing prosthesis. Therefore it is
recommended that the VA Seattle foot be re-
aligned when applying it to a prosthesis. Generally
speaking, the alignment of the VA Seattle foot is
closer to that of the SAFE and Greissinger feet
than the SACH foot. The amount of socket
flexion or plantar-flexion differs considerably
between the Seattle and SACH foot, as does the
anterior and posterior position of the foot with
respect to the socket.

As the foot is moved into plantar-flexion, the
patient will be able to notice the level of push-off
increase. However, as the level of push-off in-

FIGURE 6
Current design using Delrin® keel and Kevlar® toe
extension.
creases, this increases the hyperextension moment of the knee during midstance and considerable effort needs to be taken to walk over the foot. In alignment, plantar-flexion must be balanced with knee hyperextension. The patient will also notice the amount of push-off or spring increase as the foot is moved posterior with respect to the socket. The prosthetist should find a compromise between the hyperextension moment at midstance and the level of push-off required. The knee should not be forced into hyperextension during any phase of gait. When aligning the foot for running, the prosthetist should externally rotate the foot 2 to 3 degrees beyond its normal position, since the prosthesis has a tendency to internally rotate while running. This compensates for the rotation so both feet are symmetric.

When using the VA Seattle foot on above-knee or hip-disarticulation patient, standard multiaxis foot alignment procedures are used, except the pylon is placed in 2 to 3 degrees of posterior tilt.
### ACCEPTANCE TESTING

<table>
<thead>
<tr>
<th>SAMPLE</th>
<th>ACCEPTANCE TEST</th>
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<tbody>
<tr>
<td><strong>KEELS</strong></td>
<td>Visual Inspection for Surface Defects.</td>
</tr>
<tr>
<td></td>
<td>Weight</td>
</tr>
<tr>
<td></td>
<td>3X B.W. Load Deflection and Permanent Set.</td>
</tr>
<tr>
<td></td>
<td>Fatigue Test. Perm. Set @ Compl. Each Load Level.</td>
</tr>
<tr>
<td></td>
<td>Full Dimension Check</td>
</tr>
<tr>
<td>1 Random selected sample each configuration (beam depth and length) from 1st production batch</td>
<td></td>
</tr>
<tr>
<td>5 Random selected samples each configuration from 1st production batch.</td>
<td></td>
</tr>
<tr>
<td>1 Random sample each configuration each production batch.</td>
<td></td>
</tr>
<tr>
<td>Every 250 delivered units each configuration (units 250, 500, etc.)</td>
<td></td>
</tr>
<tr>
<td>Every 500 delivered units each configuration (units 500, 1000 etc.)</td>
<td></td>
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<tr>
<td><strong>FEET</strong></td>
<td></td>
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<tr>
<td></td>
<td>1st Production assembly each configuration.</td>
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<tr>
<td></td>
<td>1st production assembly 9 L&amp;R DS70, 10 L&amp;R DL 75, foam material batch.</td>
</tr>
<tr>
<td></td>
<td>Every foot assembly.</td>
</tr>
<tr>
<td><strong>FOAM MATERIAL</strong></td>
<td>Measure free foam density, cream and rise time, and shrinkage.</td>
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</tbody>
</table>

**FIGURE 8**
Acceptance test procedure.
FIGURE 9A
Changes in walking versus changes in running with the VA Seattle foot: test subjects' responses to whether they had noticed any changes in these 2 activities.

FIGURE 9B
Endurance: test subjects' responses to whether they could perform their sports activities for less, same, or greater amounts with this foot design.

FIGURE 9C
Improved prosthesis: test subjects' responses to whether the Seattle foot was an overall improvement compared with their previous foot.

This preloads the keel and makes the pylon vertical during midstance. If the knee becomes unstable, the prosthetist can either increase the toe lever by plantar-flexing the foot or moving the knee center posterior to the TKA line.

CLINICAL EVALUATION

In the field of mobility aids, consumer acceptance is the final measure of success. This is especially so with prostheses. Satisfaction is necessarily relative since no prosthesis fully replaces function of the missing part. Functional and psychological loss are least evident in the congenital amputee whose experience does not include the missing member. Nonetheless, all amputees are fundamentally dissatisfied to some degree with the prosthesis. This dissatisfaction cannot be overcome since the prosthesis is ultimately measured in performance against the lost body part. However, when measured against a severely deformed, painful, infected or life-threatening residual member, relief by amputa-
tion and subsequent prosthetic rehabilitation will prove a blessing. Nonetheless, the amputee must live with functional and body image alteration.

Comparisons between prostheses is another matter. Here, in most cases evaluation both subjective and objective provides a high degree of success measurement. Feedback data obtained by existing prosthetic users can be refined to provide an excellent engineering measurement tool. Physical performance such as the ability to run at certain speeds and on varied surfaces, walking endurance, and measurements of metabolic activity also add to evaluation.

Our initial evaluation of the VA Seattle foot was carried out over a 3-year period with 36 amputees. All had previously worn a prosthesis. They were selected because of interest and desire to increase physical capabilities, primarily sports. Motivation was not a problem. None were paid for participating in the program. The experimental feet, and when necessary, prosthetic adjustment were furnished without cost to the test subjects. Personal interviews and questionnaires completed by prosthetists and amputees were summarized.

The amputee acceptance rate was high. When a failure occurred, test subjects often refused to give back the failed foot for study unless a replacement was provided so as to avoid returning to the use of their former appliance. The remarkable level of acceptance was paralleled by enhanced performance, greater endurance and in most instances, comfort.

CONCLUSIONS

A prosthetic foot is described, which consists of a monolithic composite keel engineered to store potential energy and release kinetic energy (force) during ground contact. Extensive clinical evaluation indicates high user acceptance. The present design is stabilized. Technology transfer for commercial availability has been carried out.

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REFERENCES


